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DISEASE RELATED PROTEIN NETWORK

The present invention relates to a method for generating a network of direct and indirect interaction partners of a disease-related (poly)peptide comprising the steps of (a) contacting a selection of (poly)peptides suspected to contain one or several of said direct or indirect interaction partners with said disease-related (poly)peptides and optionally with known direct or indirect interaction partners of said diseaserelated (poly)peptide under conditions that allow the interaction between interaction partners to occur; (b) detecting (poly)peptides that interact with said disease-related (poly)peptide or with said known direct or indirect interaction partners of said disease-related (poly)peptide; (c) contacting (poly)peptides detected in step (b) with a selection of (poly)peptides suspected to contain one or several (poly)peptides interacting with said (poly)peptides detected in step (b) under conditions that allow the interaction between interaction partners to occur; (d) detecting proteins that interact with said (poly)peptides detected in step (b); (e) contacting said diseaserelated (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide, said (poly)peptides detected in steps (b) and (d) and a selection of proteins suspected to contain one or several (poly)peptides interacting with any of the afore mentioned (poly)peptides under conditions that allow the interaction between interaction partners to occur; (f) detecting (poly)peptides that interact with said disease-related (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide or with said (poly)peptides identified in step (b) or (d); and (g) generating a (poly)peptide -(poly)peptide interaction network of said disease-related (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide and said (poly)peptides identified in steps (b), (d) and (f). Moreover, the present invention relates to a protein complex comprising at least two proteins and to methods for identifying compounds interfering with an interaction of said proteins. Finally, the present invention relates to a pharmaceutical composition and to the use of compounds identified by the present invention for the preparation of a pharmaceutical composition for the treatment of Huntington's disease.

Several documents are cited throughout the text of this specification. The disclosure content of the documents cited herein (including any manufacture's specifications, instructions, etc.) is herewith incorporated by reference. The present invention is based on scientific experiments which have been performed on biological specimen derived from diseased patients. Patients have given their consent to use the specimen for the study which is disclosed in the present invention. In case of deceased patients, the consent has been given by a relative.

With the identification of >35.000 genes in the human genome the challenge arises to assign biological function to all proteins and to link these proteins to physiological pathways and disease processes. Since protein-protein interactions play a role in most events in a cell, clues to the function of an unknown protein can be obtained by investigating its interaction with other proteins whose function are already known. Thus, if the function of one protein is known, the function of the binding parners can be infered (deduced). This allows the researcher to assign a biological function to uncharacterized proteins by identifying protein-protein interactions. For example, several so far uncharacterized proteins in Caenorhabditis elegans were identified in a yeast two-hybrid screen for eukaryotic 26S proteasome interacting proteins and thereby could be linked to the ubiquitin-proteasome proteolytic pathway (Vidal et al., 2001). Elucidation of protein-protein interactions is particularly desired when it comes to the generation of new drugs. For many diseases, the available drug portfolio is insufficient or inappropriate to provide a cure or to prevent onset of the disease. One such disease is Huntington's disease.

Huntington's disease (HD) is a neurodegenerative disorder caused by an expanded polyglutamine (polyQ) tract in the multidomain protein huntingtin (htt). The elongated polyQ sequence is believed to confer a toxic gain of function to htt. It leads to htt aggregation primarily in neurons of the striatum and cortex and subsequently to the appearance of the disease phenotype. However, there is experimental evidence that loss of htt function may also contribute to HD pathogenesis. Since huntingtin aggregation correlates with disease progression, it is crucial to develop methods for identifying factors that promote or inhibit aggregation of huntingtin.

Previously, a number of single interaction partners of huntingtin had been reported. In light of these reports, it is tempting to speculate that huntingtin is bound into a larger network of interacting partners, many of which might be capable of modulating huntingtin's activity and function by direct or indirect interaction. It is likely that an aberrant interaction of huntingtin with some of the members of said network will impair huntingtin's normal function. Moreover, this interaction might also be relevant for the conformation of huntingtin or for its solubility or state of aggregation. Interfering with the direct or indirect interactions of the protein-protein interaction network will provide an excellent basis for therapeutic intervention as it will allow to modulate huntingtin's activity or state of aggregation or both. The state of the art so far did not provide compounds capable of reducing or suppressing huntingtin aggregation were not known.

Thus, the technical problem underlying the present invention was to provide novel approaches for identifying direct or indirect interaction partners of disease-related proteins, which must be seen as new targets for drug development. The solution to this technical problem is achieved by providing the embodiments characterized in the claims.

Accordingly, the present invention relates to a method for generating a network of direct and indirect interaction partners of a disease-related (poly)peptide comprising the steps of (a) contacting a selection of (poly)peptides suspected to contain one or several of said direct or indirect interaction partners with said disease-related (poly)peptides and optionally with known direct or indirect interaction partners of said disease-related (poly)peptide under conditions that allow the interaction between interaction partners to occur; (b) detecting (poly)peptides that interact with said disease-related (poly)peptide or with said known direct or indirect interaction partners of said disease-related (poly)peptide;(c) contacting (poly)peptides detected in step (b) with a selection of (poly)peptides suspected to contain one or several (poly)peptides interacting with said (poly)peptides detected in step (b) under conditions that allow the interaction between interaction partners to occur; (d) detecting proteins that interact with said (poly)peptides detected in step (b); (e) contacting said disease-related (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide, said (poly)peptides

detected in steps (b) and (d) and a selection of proteins suspected to contain one or several (poly)peptides interacting with any of the afore mentioned (poly)peptides under conditions that allow the interaction between interaction partners to occur; (f) detecting (poly)peptides that interact with said disease-related (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide or with said (poly)peptides identified in step (b) or (d); and (g) generating a (poly)peptide-(poly)peptide interaction network of said disease-related (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide and said (poly)peptides identified in steps (b), (d) and (f).

In accordance with the present invention, the term "direct and indirect interaction partners" relates to (poly)peptides that either directly interact with the disease-related (poly)peptide (direct interaction) or that interact via a protein binding to/interacting with said disease-related (poly)peptide. In the letter case, there is no direct contact between the direct interaction partner and the disease-related protein. Rather, a further protein forms a "bridge" between these two proteins.

The term "known direct or indirect interaction partners" refers to the fact that for certain disease-related (poly)peptides, such interaction partners are known in the art. If such interaction partners are known in the art, it is advantageous to include them into the method of the invention. If no such interactions partners are known in the art, then the network may be generated starting solely from the known disease-related (poly)peptide.

The term "conditions that allow the interaction between interaction partners to occur" relates to conditions that would, as a rule, resemble physiological conditions. Conditions that allow protein actions are well known in the art and, can be taken, for example from Golemis, E.A. Ed., Protein-Protein Interactions, Cold Spring Harbor Laboratory Press, 2002.

The term "suspected to contain one or more of said direct or indirect interaction partners" relates to the fact that normally, a selection of (poly)peptides would be employed where the person skilled in the art would expect that interaction partners

are present. Examples of such selections of (poly)peptides are libraries of human origin such as cDNA libraries or genomic libraries.

The term "detecting proteins" refers to the fact that the (poly)peptides interacting with the "bait" (poly)peptides are identified within the selection of (poly)peptides. A further characterization or isolation of the "prey" (poly)peptides at this stage may be advantageous but is not necessary. The term "detecting (poly)peptides" preferably also comprises characterizing said (poly)peptides or the nucleic acid molecules encoding said (poly)peptides. The skilled person knows that this can be done by a number of techniques, some of which are described for example in Sambrook et al., "Molecular Cloning, A Laboratory Manual"; CSH Press, Cold Spring Harbor, 1989 or Higgins and Hames (eds.). For example, the nucleotide sequence may be determined by DNA Sequencing, including PCR-Sequencing (see for example Mullis K, Faloona F, Scharf S, Saiki R, Horn G, Erlich H., Cold Spring Harb Symp Quant Biol. 1986;51 Pt 1:263-73). Alternatively, the amino acid sequence of said (poly)peptide may be determined. The skilled artesian knows various methods for sequencing proteins which include the method of Edman degradation, which is a preferred method of the present invention of determining the amino acid sequence of a protein. However, the amino acid sequence of a protein or (poly)peptide can also be reliably determined by methods such as for example Maldi-Tof, optionally in combination with the method of Edman degradation. The interaction partner may be identified either as fusion with a DNA binding domain or as fusion with an activation domain. Preferably, if an interaction partner has been identified as a fusion molecule comprising a DNA binding domain, the interaction partner is cloned into a vector allowing the expression of the interaction partner as a fusion with an activation domain. Consequently, protein interaction can be tested in the context the DNA activation or the DNA binding domain.

In accordance with the present invention, the first round of detecting (poly)peptides that interact with the "bait" (poly)peptides recited in step (a) wherein the detected (poly)peptides be considered as "prey" (poly)peptides is followed by the second round of detecting further interacting (poly)peptides wherein the former "prey" (poly)peptides are now used as "bait" (poly)peptides. In certain preferred embodiments of the present invention such as in a two-hybrid detection system, a re-

cloning of the former "prey" (poly)peptides into vectors that are suitable for expressing "bait" (poly)peptides may be desired.

Accordingly, the invention describes a novel strategy to identify protein-protein interaction networks for human disease proteins. This strategy was applied to detect pair-wise protein-protein interactions for Huntington's disease and is useful for other hereditary diseases as well. Several human hereditary diseases are summarized in table 5.

A crucial step of the method of the invention is step (e). Here, the disease-related (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide are contacted under appropriate conditions, preferably at the same time, with both the (poly)peptides identified in steps (b) and (d) and further with a selection of (poly)peptides suspected to contain further interaction partners. Alternatively, the various baits, preys and further selection partners are added one after another, so that the final pool contains all baits and preys so far identified, in addition to the further selection partners. In other terms, in this step of the method of the invention, all "baits" and all "preys" are pooled and, additionally, further potential interaction partners are added. In this way, surprisingly the number of directed or indirect interactions partners of the previously identified "baits" and "preys" could significantly be enhanced. It is to be understood that various preys identified in one detection step may interact with each other and not only with the baits that were employed for the identification. For example, if a collection of baits detects prays "a" and "b", the invention does not exclude that "a" also interacts with "b". The same holds true mutatis mutandis for the baits used in accordance with the present invention. Wherever possible, baits and preys are exchangeable in the sense that bait (poly)peptides may be used as preys and vice versa. In a given case, however, the skilled person has to determine whether or not this exchange is possible on the basis of unfavourable site effects and limitations of the applied scientific approach. This can be done by the skilled person without undue burden by applying standard techniques known in the art.

It is further preferred in accordance with the present invention that the interaction of proteins is a specific interaction, such as a specific binding. This means that the

(poly)peptide being an interaction partner with a further (poly)peptide only or essentially only interacts with the interaction site(s) involved with this interaction partner. This does not exclude, of course, that further interaction sites of said (poly)peptide interact with further interaction partners, wherein in the corresponding interaction is preferably also specific. The concept also embraces that, if a (poly)peptide has several identical interaction sites, which in nature bind to different interaction partners, these different interaction partners are also bound by the (poly)peptide in the method of the present invention.

In other terms, at least in the case of huntingtin, the number of interaction partners found in step (e) was enhanced in an exponential rather than in a linear fashion.

The term "(poly)peptide" refers alternatively to peptide or to (poly)peptides. Peptides conventionally are covalently linked amino acids of up to 30 residues, whereas polypeptides (also referred to as "proteins") comprise 31 and more amino acid residues.

The term "huntingtin" refers to a protein with the data bank accession number P42858 which is referenced for the purpose of the present invention as "wild-type huntingtin protein". However, the term "huntingtin" also comprises proteins encoded by the nucleic acid sequence deposited under accession number L12392 or to proteins encoded by nucleic acid molecules which hybridize to the nucleic acid molecule of L12392 under stringent conditions of hybridization. The present invention relates to all variants of the huntingtin protein. In particular, relevant for the present invention are those variants of huntingtin which comprise a polyglutamine tract (polyQ troct) or an elongated polyQ tract. A polyQ tract consists of two or more glutamines within the huntingtin protein. The insertion of additional glutamine codons will result in huntingtin proteins with, for example 2, 51, 75 or 100 added glutamines in comparison to the sequence deposited under accession number P42858. In fact, the person skilled in the art knows that the huntingtin protein may have a glutamine tract with any random number of glutamines in the range of 1 to 200 added glutamines. All these proteins are comprised by the present invention.

The term "hybridizes under stringent conditions", as used in the description of the present invention, is well known to the skilled artisian and corresponds to conditions

of high stringency. Appropriate stringent hybridization conditions for each sequence may be established by a person skilled in the art on well-known parameters such as temperature, composition of the nucleic acid molecules, salt conditions etc.; see, for example, Sambrook et al., "Molecular Cloning, A Laboratory Manual"; CSH Press, Cold Spring Harbor, 1989 or Higgins and Hames (eds.), "Nucleic acid hybridization, a practical approach", IRL Press, Oxford 1985, see in particular the chapter "Hybridization Strategy" by Britten & Davidson, 3 to 15. Stringent hybridization conditions are, for example, conditions comprising overnight incubation at 42° C in a solution comprising: 50% formamide, 5x SSC (750 mM NaCl, 75 mM trisodium citrate), 50 mM sodium phosphate (pH 7.6), 5x Denhardt's solution, 10% dextran sulfate, and 20 micrograms/ml denatured, sheared salmon sperm DNA, followed by washing the filters in 0.1x SSC at about 65°. Other stringent hybridization conditions are for example 0.2 x SSC (0.03 M NaCl, 0.003M Natriumcitrat, pH 7) bei 65°C. In addition, to achieve even lower stringency, washes performed following stringent hybridization can be done at higher salt concentrations (e.g. 5X SSC). Note that variations in the above conditions may be accomplished through the inclusion and/or substitution of alternate blocking reagents used to suppress background in hybridization experiments. Typical blocking reagents include Denhardt's reagent, BLOTTO, heparin, denatured salmon sperm DNA, and commercially available proprietary formulations. The inclusion of specific blocking reagents may require modification of the hybridization conditions described above, due to problems with compatibility.

The skilled person knows that the presence of additional codons in the nucleic acid sequence of huntingtin might significantly reduce the capability of this nucleic acid molecule to hybridize to the nucleic acid molecule deposited under L12392 and referenced as wild-type huntingtin protein. Nevertheless, such proteins shall still be comprised by the present invention. In fact, computer programs such as the computer program Bestfit (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, 575 Science Drive, Madison, WI 53711) or blast, capable of calculating homologies between two nucleic acid sequences, efficiently recognize nucleotide insertions and allow for an adjustment of gaps created by these insertions. The term "huntingtin" as used in the present invention, also includes those molecules of huntingtin, which have a

homology of more than 95% to wild-type huntingtin when analyzed with a program like bestfit under conditions not weighing gaps created by polyQ tracts (gap penalty=0).

The term "contacting" means bringing into contact so that two or more proteins or (poly)peptides can interact with each other, preferably under physiological conditions. The terms "interacting" or "binding" refer to a transient or permanent contact between two proteins or (poly)peptides. Preferably, the (poly)peptide or protein is provided by expression from a nucleic acid molecule, more preferably from a cDNA molecule within a cDNA library. Alternatively, said nucleic acid molecule is a genomic nucleic acid molecule of a genomic DNA library, or a nucleic acid molecule from a synthetic DNA or RNA library. Preferably, the nucleic acid molecule encoding the diseaserelated protein or its interaction partner is obtainable from nerve cells, brain tissue human adrenal gland, human bladder, human bone, human brain, human colon, human dorsal root ganglion, human heart, human HeLa cells, human kidney, human liver, human lung, human mammary gland, human ovary, human pancreas, human placenta, human prostate, human retina, human salivary gland, human sceletal muscle, human small intestine, human smooth muscle, human spinal cord, human spleen, human stomach, human testis, human thymus, human thyroid, human tonsil, human trachea, human uterus, human cell line HEP G2, human cell line MDA 435, human fetal brain, human fetal heart, human fetal kidney, human fetal liver, human fetal spleen, human fetal thymus, human breast tumor, human cervix tumor, human colon tumor, human kidney tumor, human lung tumor, human ovary tumor, human stomach tumor, human brain tumor and/or human uterus tumor.

The term "disease-related protein" refers to a protein known to be the causative agent of a disease or known to be involved in onset or progression of a disease. Preferably, said disease is CHOREA HUNTINGTON or the disease-related protein is huntingtin. More preferably, the disease-related protein is selected from table 6 and/or 7. The term "conditions that allow the interaction between interaction partners" means conditions that are similar to physiological conditions. Preferably, said conditions are physiological conditions.

The term "selection of (poly)peptides" refers to a library of (poly)peptides which comprises the above-mentioned libraries, but also includes libraries such as phage

display libraries. Preferably, the (poly)peptide is provided by expression from a nucleic acid molecule. Preferably, the protein or (poly)peptide expressed by said nucleic acid molecule is a (poly)peptide comprising a DNA binding domain (DBD) (in this case the fusion protein is termed "bait") or (b) a (poly)peptide comprising an activation domain capable of interacting with a transcription factor or an RNA polymerase and capable of activating transcription of a reporter or indicator gene (in this case the fusion protein is called "prey"). As used here, the terms "reporter gene" and "indicator gene" are to be understood as synonyms. It is important to note that one of the interaction partners will always comprise the amino acid sequence of a protein or (poly)peptide translated from said nucleic acid molecule while the other interaction partner will comprise the amino acid sequence of a protein or protein fragment. Preferably, a bait used for a method of the present invention is selected from the proteins listed in table 6 and/or 7. If, for example, the proteins encoded by the nucleic acid molecules contain a DNA binding domain fused in frame, the fusion protein can bind to the DNA recognition sequence of the DNA binding domain. Interaction of said fusion protein with a second fusion protein containing an activation domain can induce transcription of a nearby indicator gene. The indicator gene may encode a selection marker such as a protein that confers resistance to an antibiotic including ampicillin, kanamycin, chloramphenicol, tetracyclin, hygromycin, neomycin or methotrexate. Further examples of antibiotics are Penicillins: Ampicillin HCl, Ampicillin Na, Amoxycillin Na, Carbenicillin disodium, Penicillin G, Cephalosporins, Cefotaxim Na, Cefalexin HCl, Vancomycin, Cycloserine. Other examples include Bacteriostatic Inhibitors such as: Chloramphenicol, Erythromycin, Lincomycin, Tetracyclin, Spectinomycin sulfate, Clindamycin HCl, Chlortetracycline HCl. Additional examples are proteins that allow selection with Bacteriosidal inhibitors such as those affecting protein synthesis irreversibly causing cell death. Aminoglycosides can be inactivated by enzymes such as NPT II which phosphorylates 3'-OH present on kanamycin, thus inactivating this antibiotic. Some aminoglycoside modifying enzymes acetylate the compound and block their entry in to the cell. Gentamycin, Hygromycin B, Kanamycin, Neomycin, Streptomycin, G418, Tobramycin Nucleic Acid Metabolism Inhibitors, Rifampicin, Mitomycin C, Nalidixic acid, Doxorubicin HCI, 5-Flurouracil, 6-Mercaptopurine, Antimetabolites, Miconazole, Trimethoprim, Methotrexate, Metronidazole, Sulfametoxazole. Alternatively, said indicator gene may encode a protein such as lacZ, GFP or luciferase, the expression

of which can be monitored by detection of a specific color. Other proteins commonly used as indicator proteins are beta-galactosidase, beta-glucuronidase, green fluorescent protein (GFP), autofluorescent proteins, including blue fluorescent protein (BFP), glutathione-S-transferase (GST), luciferase, horseradish peroxidase (HRP), and chloramphenicol acetyltransferase (CAT). In general, however, the selection in the yeast two hybrid-system is based on a deficiency of the yeast strain to produce specific amino acids. The skilled person knows that any amino acid deficiency can be used for this selection strategy.

Preferably said preys and baits are expressed from two separate expression vectors contained in one host cell. The nucleic acid molecule encoding the preys and baits can be introduced into the host cell, for example, by transformation, transfection, transduction or microinjection which are common techniques known to the person skilled in the art and which require no additional explanation. In addition, the nucleic acid molecule contains a chromosomal or episomal nucleic acid sequence encoding the above-mentioned indicator protein. The expression of said indicator protein is under control of a recognition sequence which serves as a binding site for the bait protein. The nucleic acid molecule may be fused either to a DNA binding domain or to an activation domain. Co-expression of only those bait- and prey fusion proteins which are capable of interacting will induce the expression of one of the aboveidentified indicator proteins and thus allow the identification a nucleic acid molecule encoding a protein capable of interacting with huntingtin or an interaction or binding partner of huntingtin. The skilled person knows this system as the yeast two hybrid system. The yeast two hybrid system, which uses a bait protein-prey protein combination to induce transcription of the reporter gene, is a preferred method to identify proteins capable of interacting with huntingtin or with a direct or indirect interaction or binding partner of huntingtin. See for example Fields and Song, Nature 340:245 (1989) or Uetz et al., 2000 Nature 403(6770): 623-7. This is a useful way of determining protein-protein interactions. Another preferred method uses the yeast three hybrid system, as described in U.S. Pat. No. 5,928,868. Preferably, steps (a) to (d) of the method for generating a network of direct and indirect interaction partners comprise the yeast two hybrid system. Preferably, steps (e) and (f) of the method for generating a network of direct and indirect interaction partners comprise yeast interaction mating. Preferably, said "interaction mating" comprises the interaction of all interaction partners identified in steps (a) to (d). Also preferred is that the interaction partners identified in steps (a) to (d) interact as prey and bait proteins, so that all prey proteins are contacted with all bait proteins. Using the array mating system, each bait is tested individually for interaction with every prey in the array. Alternatively, steps (e) and (f) of the method for generating a network of direct and indirect interaction partners comprise testing all interaction partners identified in steps (a) to (d) in interaction assays such as biacore or coimmunoprecipitation. When performing such an assay, it is preferred that the interaction partners are tested as prey and/or bait fusion proteins or contain no fused (poly)peptides. Preferably, all interaction partners are contacted in the biacore or coimmunoprecipitation assay by themselves and by all other remaining interaction partners identified in steps (a) to (d).

The method for generating a network of direct and indirect interaction partners of a disease related protein or (poly)peptide has proven to be an effective tool for unveiling the protein-protein interactions (PPI) of preferably monogenic diseases. This is exemplified by the analysis of the disease related protein of Chorea Huntington, the analysis of which has demonstrated that the method of the present invention will be useful in an approach to identify potential drugs in the treatment of CHOREA HUNTINGTON. Moreover, this method will also be effective in unveiling the protein-protein interactions of other disease related proteins and in identifying novel targets for treatment of these diseases. Using a preferred combination of library and matrix yeast two-hybrid screens, based on the methods of the present invention, a highly connected network was generated among 70 proteins involved in 117 protoin-protein interactions, 99 of which had not been described previously. As progression of Huntington's disease (HD) appears to be linked to huntingtin aggregation, a set of network proteins was tested for their potential to modulate this process. By using the methods of the present invention, it was discovered that the GTPase activating protein GIT1 strongly promotes huntingtin aggregation in vivo. GIT1 also localises to huntingtin aggregates in brains of transgenic mice and HD patients. Therefore, a combination of the methods of the present invention has proven to provide effective means for the identification of potential targets for therapeutic intervention. GIT1 is a selected example of a modulator interaction partner of huntingtin. The other proteins in the network of interaction partners

disclosed by the present invention are further modulator interaction partners of huntingtin.

Preferably, the interaction mating comprises using an array maiting system. In general, for this screen, MATa yeast cultures are transformed with plasmids encoding prey proteins and arrayed on a microtiter plate for interaction mating with individual MATa strains expressing bait proteins. Using this test system, each bait can be tested individually for interaction with every prey in the array. Diploid yeast clones, formed by maiting on YPD plates and expressing both, bait and prey proteins, are selected on agar SDII plates, and further transferred for example by a spotting robot on SDIV plates to select for protein-protein interactions. In a more preferred embodiment of the method, plasmids encoding bait and prey proteins are transformed into strains L40ccua and L40cca, respectively. L40cca clones are arrayed on microtitre plates and mixed with a single L40ccua clone for interaction mating. These cells are transferred, preferably by a robot onto YPD medium plates and, after incubation for 20h to 28h at approximately 30°C, for selection of the cells, were transferred onto SDII medium plates, where mating takes place, for additional 60h to 80h at approximately 30°C. For two-hybrid selection diploid cells are transferred onto SDIV medium plates with and without nylon or nitrocellulose membranes and incubated for approximately 5 days at about 30°C. The nylon or nitrocellulose membranes are subjected to the β-GAL assay. Positive clones can be verified by cotransformation assays using plasmids encoding respective bait and prey proteins. Other preferred methods for studying protein-protein interactions according to the present invention are colocalization, coimmunoprecipitation, screening of protein or (poly)peptide arrays, library screens, in vivo and in vitro binding experiments using different tags such as HIS6, TAP or FLAG.

In a preferred embodiment of the present invention's method for generating a network of direct and indirect interaction partners of a disease related protein or (poly)peptide, plasmids encoding bait proteins are transformed into a strain such as L40ccua, tested for the absence of reporter gene activity and co-transformed with a human fetal brain cDNA library. Independent transformants are plated onto minimal medium lacking tryptophan, leucine, histidine and uracil (SDIV medium) and incubated at about 30°C for 5 to 10 days. Clones are transferred into microtitre

plates, optionally using a picking robot, and grown over night in liquid minimal medium lacking tryptophan and leucine (SDII medium). Subsequently, the clones are spotted onto nylon or nitrocellulose membranes placed on SDIV medium plates. After incubation for about 4 days membranes are subjected to a β -galactosidase (β -GAL) assay. Plasmids are prepared from positive clones and characterised, for example by restriction analyses and sequencing. For retransformation assays plasmids encoding bait and prey proteins are cotransformed in the yeast strain L40ccua and plated onto SDIV medium.

The term "generating a protein-protein interaction (PPI) network" means listing the interactions of all proteins interacting or binding directly or indirectly interacting the disease related (poly)peptide or protein. Preferably, this can be done by displaying the information in a matrix or a network representation. In a more preferred embodiment of the present invention's method, the protein-protein interaction network is generated by using Pivot 1.0 (Prof. Ron Shamir, Prof. Yossi Shilo, Nir Orlev; Tel Aviv University (TAU); Dep. of computer science; Ramat Aviv; Tel Aviv 69978; Israel).

In a preferred embodiment of the invention, interactions are detected by using the yeast two-hybrid system, MALDI-TOF MS or electro spray MS. Preferably, yeast strains such as strains L40ccua and L40cca, are transformed with an expression selected from the group consisting of pBTM116, pBTM117, pBTM117c, pACT2, pAS2-1, pGAD10, pGAD424, pGAD425, pGAD426, pGAD427, pGAD428.

In another preferred embodiment of the present invention's method for generating a network of direct and indirect interaction partners of a disease-related polypeptide, the method contains after step (d) the additional steps of isolating a nucleic acid molecule with homology to said nucleic acid molecule expressing the encoded protein and testing it for its activity as a modulator of huntingtin, wherein said nucleic acid molecule is DNA, RNA, cDNA, or genomic DNA. Said testing can be done in several different assays. Preferably, the testing is performed in a co-immunoprecipitation assay or an affinity chromatography-based technique. Generally, co-immunoprecipitation is performed by purifying an interacting protein complex with a single antibody specific for one protein in the protein complex and by detecting the proteins in the protein complex. The step of detection can involve the

use of additional antibodies directed against proteins suspected of being trapped in the purified protein complex. Alternatively, at least one protein in the protein complex is fused to a tag sequence with affinity to a compound fixed to a solid matrix. By contacting the solid matrix with said tagged protein, further proteins binding to said protein can be purified and binding can be detected. GST or HA are preferred tags in accordance with the present invention.

In a preferred embodiment of the present invention's method, said contacting step (e) is effected in an interaction mating two hybrid approach.

In another preferred embodiment of the present invention's method, said method comprises after step (d) and before step (e) the steps of: (d') contacting (poly)peptides detected in step (d) with a selection of (poly)peptides suspected to contain one or several (poly)peptides interacting with said (poly)peptides detected in step (d) under conditions that allow the interaction between interaction partners to occur; and (d") detecting proteins that interact with said (poly)peptides detected in step (d').

This preferred embodiment of the invention, an additional step of identifying further interaction partners is carried out prior to the contacting of all "baits" and "preys" in one pool (step (e)). Optionally, further steps of selecting interaction partners in analogy to steps (d') and (d") may be infected prior to the pooling/interaction step.

Diseases of particular interest for which interrelationships of disease-related proteins may be analyzed in accordance with the invention are provided in Table 5.

In yet another preferred embodiment of the present invention's method, said disease related protein is a protein suspected of being a causative agent of a hereditary (see Table 5), such as a monogenic disease.

In another preferred embodiment of the present invention's method, said disease related protein is huntingtin and said interaction partners are the interaction partners as shown in table 6,7 and/or 9

In another preferred embodiment of the present invention's method, said method comprises the step of determining the nucleotide sequence of a nucleic acid molecule encoding a direct or indirect interaction partner of the disease related

In another preferred embodiment of the present invention's method, said selections of proteins are translated from a nucleic acid library.

protein.

In another preferred embodiment of the present invention's method, said selection of proteins in step (a) and/or (c) and/or (d') and/or (e) is the same selection or a selection from the same source. In another preferred embodiment of the present invention's method, said selection of proteins in step (a) and/or (c) and/or (d') and/or (e) is a different selection or a selection from a different source.

Preferably, said source is selected from nerve cells, brain tissue, human adrenal gland, human bladder, human bone, human brain, human colon, human dorsal root ganglion, human heart, human HeLa cells, human kidney, human liver, human lung, human mammary gland, human ovary, human pancreas, human placenta, human prostate, human retina, human salivary gland, human sceletal muscle, human small intestine, human smooth muscle, human spinal cord, human spleen, human stomach, human testis, human thymus, human thyroid, human tonsil, human trachea, human uterus, human cell line HEP G2, human cell line MDA 435, human fetal brain, human fetal heart, human fetal kidney, human fetal liver, human fetal spleen, human fetal thymus, human breast tumor, human cervix tumor, human colon tumor, human kidney tumor, human lung tumor, human ovary tumor, human stomach tumor, human brain tumor and/or human uterus tumor.

In another preferred embodiment of the present invention's method, said method is performed by contacting the proteins on an array. Preferably, said array is an array allowing to detect protein-protein interaction by the principle of a biacore detector.

In another preferred embodiment of the present invention's method, said interactions are detected by using the yeast two-hybrid system. Preferably, said interactions detected by using MALDI-TOF, MS, electro spray MS or biacore.

In another preferred embodiment of the present invention's method, said method contains after step of (b), (d), (d") or (f) the additional steps of isolating a nucleic acid molecule with homology to said cDNA expressing the encoded protein and testing it for its activity as a modulator of huntingtin, wherein said nucleic acid molecule is DNA, or RNA, and preferably cDNA, or genomic or synthetic DNA, or mRNA.

By using the methods disclosed herein, a rate of success or fidelity of at least 70% validatable protein-protein interactions (PPI) (of proteins within the protein interaction network of huntingtin) can be achieved. This level of consistency is well above the level described in the art. In order to increase the rate of success or fidelity, the skilled person can, when carrying out the methods of the present invention, combine the methods of the present invention with additional steps of testing. For example, a step of co-immunoprecipitation and/or an in vitro binding assay may be carried out, in cases when initially the interaction was determined by using the yeast-two-hybrid system (or vice versa). Such additional steps may be carried out at any stage of the methods of the present invention. For example, after but also prior to step (f) of the method of the present invention, PPIs may be verified using in-vitro binding and/or immunoprecipatation assays in order to increase the stringency of the method. By performing these additional steps of testing, the skilled person can increase the rate of success or fidelity to at least 50%, more preferably to at least 60%. For the additional validation, any method may be employed that is available to the skilled artisan for testing the protein interaction. For example, the skilled artisan may simply repeat the step(s) initially carried out, optionally by (slightly) altering the reaction conditions, preferably to more stringent reaction conditions, i.e. conditions that could be expected to further reduce the number of false positive interactions. Alternatively, a different method may be carried out in the validation process. For example, if the method of the invention employed two hybrid systems, the validation might be carried out by precipitation steps as outlined elsewhere in the specification. Whereas the method of the invention provides valid results without the additional validation step(s), the inclusion of such additional validation steps may be advantageous for certain purposes, e.g. drug target identification. In the case that a first validation step does not confirm that the protein in question is a member of the interaction network, further steps in this regard should be carried out. For example, it should be excluded that the validation step(s) do/does not catch weak protein interactions that

nevertheless are part of the network. The present invention also relates to a nucleic acid molecule encoding a modulator of huntingtin, wherein said modulator is a protein selected from table 8. Figure 6 provides the amino acid sequences of the new proteins or (poly)peptides listed in table 8. The term "modulator protein of huntingtin" comprises two types of proteins within the network of proteins interacting with huntingtin. Direct interaction or binding partners of huntingtin are those proteins in the PPI network of huntingtin that directly interact with or bind to huntingtin (see figure 2). Examples of these proteins are IKAP, HYPA, CA150, HIP1, HIP11, HIP13, HIP15, CGI-125, PFN2, HP28, DRP-1, SH3GL3, HZFH, HIP5, PIASy, HIP16, GIT1, Ku70 and FEZ1. Table 7 and figure 6 provides a reference allowing to identify these proteins. The second class of proteins are indirect interaction or binding partners of huntingtin, i.e. those proteins in the PPI network of huntingtin that do not directly interact with or bind to huntingtin. Such proteins require a mediator, i.e. a direct binding partner of huntingtin to exert their huntingtin modulating function. Examples of these proteins are BARD1 or VIM, which bind to direct interaction partners of huntingtin. However, complexes of huntingtin and a direct interaction or binding partner are likely to interact with additional indirect interaction or binding partners. To summarize the above, modulator proteins of huntingtin can exert their function by direct or indirect contact to huntingtin.

The term "modulator protein", as used in the present invention, refers to a protein capable of modulating the function or physical state of a second protein and comprises proteins that enhance or reduce (inhibit) the function or activity of huntingtin. Preferably, the modulator protein is a protein having an activity selected from the group consisting of oxidoreductase activity (acting on the CH-OH group of donors, acting on the aldehyde or oxo group of donors, acting on the CH-CH group of donors, acting on the CH-NH(2) group of donors, acting on the CH-NH group of donors, acting on NADH or NADPH, acting on other nitrogenous compounds as donors, acting on a sulfur group of donors, acting on a heme group of donors, acting on diphenols and related substances as donors, acting on a peroxide as acceptor, acting on hydrogen as donor, acting on single donors with incorporation of molecular oxygen, acting on the CH-OH group of donors, acting on superoxide as acceptor, oxidizing metal ions, acting on -CH(2) groups, acting on iron-sulfur proteins as donors, acting on reduced flavodoxin as donor, acting on phosphorus or arsenic in

donors, acting on x-H and y-H to form an x-y bond, other oxidoreductases), transferase activity (transferring one-carbon groups, transferring aldehyde or ketone residues, acyltransferases, glycosyltransferases, transferring alkyl or aryl groups, other than methyl groups, transferring nitrogenous groups, transferring phosphorouscontaining groups, transferring sulfur-containing groups, transferring seleniumcontaining groups), hydrolase activity (glycosylase activity, acting on ether bonds, acting on peptide bonds, acting on carbon-nitrogen bonds (other than peptide bonds), acting on acid anhydrides, acting on carbon-carbon bonds, acting on halide bonds, acting on phosphorus-nitrogen bonds, acting on sulfur-nitrogen bonds, acting on carbon-phosphorus bonds, acting on sulfur-nitrogen bonds, acting on carbonphosphorus bonds, acting on sulfur-sulfur bonds, acting on carbon-sulfur bonds, lyases (carbon-carbon lyases, carbon-oxygen lyases, carbon-nitrogen lyases, carbon-sulfur lyases, carbon-halide lyases, phosphorus-oxygen lyases, other lyases), isomerases (racemases and epimerases, cis-trans-isomerases, intramolecular lyases, oxidoreductases, intramolecular transferases, intramolecular isomerases), ligases activity (forming carbon-oxygen bonds, forming carbon-sulfur bonds, forming carbon-nitrogen bonds, forming carbon-carbon bonds, forming phosphoric ester bonds), transcription factor activity, filament protein, membrane protein and structural protein.

In a preferred embodiment, the present invention's nucleic acid molecule is DNA, or RNA, and preferably cDNA, or genomic DNA or synthetic DNA or mRNA

In another preferred embodiment of the invention, the nucleic acid molecule is double stranded or single stranded.

In another preferred embodiment of the invention, the nucleic acid molecule is of vertebrate, nematode, insect, bakterium or yeast. Preferably, the nematode is Caenorhabditis elegans. In another more preferred embodiment of the present invention, the insect is drosophila, preferably drosiphila melanogaster. In another more preferred embodiment of the present invention, the vertebrate is human, mouse rat, Xenopus laevis, zebrafish.

In yet another preferred embodiment of the present invention, the nucleic acid molecule is fused to a heterologous nucleic acid molecule. In a further preferred

embodiment of the present invention, the heterologous (poly)peptide encoded by said heterlogous nucleic acid molecule is an immunoglobulin Fc domain.

In another preferred embodiment of the present invention the nucleic acid molecule is labeled. Labeled nucleic acid molecules may be useful for purification or detection. Suitable labels include fluorochromes, e.g. fluorescein isothiocyanate (FITC), rhodamine, Texas Red, phycoerythrin, allophycocyanin, 6-carboxyfluorescein (6-2',7'-dimethoxy-4',5'-dichloro-6-carboxyfluorescein (JOE), 6-carboxy-X-FAM). 6-carboxy-2',4',7',4,7-hexachlorofluorescein (HEX). rhodamine(ROX), carboxyfluorescein (5-FAM) or N,N,N',N'-tetramethyl-6-carboxyrhodamine (TAMRA), radioactive labels, e.g. ³²P, ³⁵S, ³H; etc. The label may also be a two stage system, where the DNA is conjugated to biotin, haptens, etc. having a high affinity binding partner, e.g. avidin, specific antibodies, etc., where the binding partner is conjugated to a detectable label. In the case of amplification the label may be conjugated to one or both of the primers. The pool of nucleotides used in the amplification may also be labeled, so as to incorporate the label into the amplification product. Alternatively, the double strand formed after hybridization can be detected by anti-double strand DNA specific antibodies or aptamers etc.

In a more preferred embodiment said heterologous nucleic acid molecule encodes a heterologous polypeptide. Preferably said heterologous (poly)peptide, fused to the (poly)peptide encoded by the nucleic acid molecule of the present invention, is a DNA binding protein selected from the group consisting of GAL4 (DBP) and LexA (DBP). Also preferred in accordance with the present invention are activation domains selected from the group consisting of GAL4(AD) and VP16(AD). Also preferred are (poly)peptides selected from the group consisting of GST, His Tag, Flag Tag, Tap Tag, HA Tag and Protein A Tag.

Thus, the sequence encoding the (poly)peptide may be fused to a marker sequence, such as a sequence encoding a peptide which facilitates purification of the fused (poly)peptide. In certain preferred embodiments of this aspect of the invention, the marker amino acid sequence is a hexa-histidine peptide, such as the tag provided in a pQE vector (QIAGEN, Inc., 9259 Eton Avenue, Chatsworth, CA, 91311), among others, many of which are commercially available. As described in Gentz et al., Proc. Natl. Acad. Sci. USA 86:821-824 (1989), for instance, hexa-histidine provides for

convenient purification of the fusion protein. The "HA" tag is another peptide useful for purification which corresponds to an epitope derived from the influenza hemagglutinin protein, which has been described by Wilson et al., Cell 37: 767 (1984).

The (poly)peptide may be expressed in a modified form, such as a fusion protein, and may include not only secretion signals, but also additional heterologous functional regions. For instance, a region of additional amino acids, particularly charged amino acids, may be added to the N-terminus of the (poly)peptide to improve stability and persistence in the host cell, during purification, or during subsequent handling and storage. Also, peptide moieties may be added to the (poly)peptide to facilitate purification. Such regions may be removed prior to final preparation of the (poly)peptide. The addition of peptide moieties to (poly)peptides to engender secretion or excretion, to improve stability and to facilitate purification, among others, are familiar and routine techniques in the art. A preferred fusion protein comprises a heterologous region from immunoglobulin that is useful to stabilize and purify proteins.

The present invention also relates to a method of producing a vector comprising the nucleic acid molecule the present invention. Furthermore, the present invention relates to a vector produced said method.

The present invention also relates to a vector comprising the nucleic acid molecule of the present invention. Preferably said vector is a transfer or expression vector selected from the group consisting of pACT2; pAS2-1; pBTM116; pBTM117; pcDNA3.1; pcDNAI; pECFP; pECFP-C1; pECFP-N1; pECFP-N2; pECFP-N3; pEYFP-C1; pFLAG-CMV-5 a, b, c; pGAD10; pGAD424; pGAD425; pGAD427; pGAD428; pGBT9; pGEX-3X1; pGEX-5X1; pGEX-6P1; pGFP; pQE30; pQE30N; pQE30-NST; pQE31; pQE31N; pQE32; pQE32N; pQE60; pSE111; pSG5; pTET-CMV-AS; pTET-CMV-F°-AS; pTET-CMV-F°-S; pTET-CMV-MCS; pTET-CMV-S; pTK-Hyg; pTL1; pTL10; pTL-HA0; pTL-HA1; pTL-HA2; pTL-HA3; pBTM118c; pGEX-6P3; pACGHLT-C; pACGHLT-A; pACGHLT-B; pUP; pcDNA3.1-V5His; pMalc2x. Said expression vectors may particularly be plasmids, cosmids, viruses or bacteriophages used conventionally in genetic engineering plasmids, cosmids, viruses and bacteriophages used conventionally in genetic engineering that comprise the

aforementioned nucleic acid. Preferably, said vector is a gene transfer or targeting vector. Expression vectors derived from viruses such as retroviruses, vaccinia virus, adeno-associated virus, herpes viruses, or bovine papilloma virus, may be used for delivery of the nucleic acid into targeted cell population. Methods which are well known to those skilled in the art can be used to construct recombinant viral vectors; see, for example, the techniques described in Sambrook et al., Molecular Cloning A Laboratory Manual, Cold Spring Harbor Laboratory (1989) N.Y. and Ausubel et al., Current Protocols in Molecular Biology, Green Publishing Associates and Wiley Interscience, N.Y. (1989).

In yet a further preferred embodiment of the invention the vector contains an additional expression cassette for a reporter protein, selected from the group consisting of ß-galactosidase, luciferase, green fluorescent protein and variants thereof.

Preferably, said vector comprises regulatory elements for expression of said nucleic acid molecule. Consequently, the nucleic acid of the invention may be operatively linked to expression control sequences allowing expression in eukaryotic cells. Expression of said nucleic acid molecule comprises transcription of the sequence nucleic acid molecule into a translatable mRNA. Regulatory elements ensuring expression in eukaryotic cells, preferably mammalian cells, are well known to those skilled in the art. They usually comprise regulatory sequences ensuring initiation of transcription and, optionally, a poly-A signal ensuring termination of transcription and stabilization of the transcript, and/or an intron further enhancing expression of said nucleic acid. Additional regulatory elements may include transcriptional as well as translational enhancers, and/or naturally-associated or heterologous promoter regions. Possible regulatory elements permitting expression in eukaryotic host cells are the AOX1 or GAL1 promoter in yeast or the CMV-, SV40-, RSV-promoter (Rous sarcoma virus), CMV-enhancer, SV40-enhancer or a globin intron in mammalian and other animal cells. Beside elements which are responsible for the initiation of transcription such regulatory elements may also comprise transcription termination signals, such as the SV40-poly-A site or the tk-poly-A site, downstream of the nucleic acid molecule. Furthermore, depending on the expression system used leader sequences capable of directing the (poly)peptide to a cellular compartment or secreting it into the medium may be added to the coding sequence of the aforementioned nucleic acid and are well known in the art. The leader sequence(s) is (are) assembled in appropriate phase with translation, initiation and termination sequences, and preferably, a leader sequence capable of directing secretion of translated protein, or a portion thereof, into the periplasmic space or extracellular medium. Optionally, the heterologous sequence can encode a fusion protein including an C- or N-terminal identification peptide imparting desired characteristics, e.g., stabilization or simplified purification of expressed recombinant product. In this context, suitable expression vectors are known in the art such as Okayama-Berg cDNA expression vector pcDVI (Pharmacia), pCDM8, pRc/CMV, pcDNA1, pcDNA3, the EchoTM Cloning System (Invitrogen), pSPORT1 (GIBCO BRL) or pRevTet-On/pRevTet-Off or pCI (Promega).

The present invention also relates to a method of producing a host cell comprising genetically engineering cells with the nucleic acid molecule or the vector of the present invention. The present invention also relates to a host cell produced said method. Furthermore, the present invention relates to a host cell comprising the vector of the present invention. Preferably, said host cell contains an endogenous nucleic acid molecule which is operably associated with a heterologous regulatory control sequence, including the regulatory elements contained in the vector of the present invention.

The present invention also relates to a method of producing a (poly)peptide, comprising culturing the host cell of the present invention under conditions such that the (poly)peptide encoded by said polynucleotide is expressed and recovering said (poly)peptide.

The present invention also relates to a (poly)peptide comprising an amino acid sequence encoded by a nucleic acid molecule of the present invention, or which is chemically synthesized, or is obtainable from the host cell of the present invention, or which is obtainable by a method of the present invention or which is obtainable from an in vitro translation system by expressing the nucleic acid molecule of the present invention or the vector of the present invention.

In another preferred embodiment of the invention, the (poly)peptide or protein is of vertebrate, nematode, insect, bakterium or yeast. Preferably, the nematode is Caenorhabditis elegans. In another more preferred embodiment of the present invention, the insect is Drosophila, preferably Drosophila melanogaster. In another more preferred embodiment of the present invention, the vertebrate is human, mouse rat, Xenopus laevis, zebrafish.

In another preferred embodiment, the (poly)peptide of the present invention is fused to a heterologous (poly)peptide. Such a fusion protein may include not only secretion signals, but also additional heterologous functional regions. For instance, a region of additional amino acids, particularly charged amino acids, may be added to the N-terminus of the (poly)peptide to improve stability and persistence in the host cell, during purification, or during subsequent handling and storage. Also, peptide moieties may be added to the (poly)peptide to facilitate purification. Such regions may be removed prior to final preparation of the (poly)peptide. The addition of peptide moieties to (poly)peptides to engender secretion or excretion, to improve stability and to facilitate purification, among others, are familiar and routine techniques in the art. A preferred fusion protein comprises a heterologous region from immunoglobulin that is useful to stabilize and purify proteins.

In a preferred embodiment of the present invention, the (poly)peptide of the present invention is fused to a heterologous (poly)peptide which is an immunoglobulin Fc domain or Protein A domain. In another preferred embodiment of the present invention, the (poly)peptide the (poly)peptide is labelled. Preferably, the label is selected from the group consisting of fluorochromes, e.g. fluorescein isothiocyanate (FITC), rhodamine, Texas Red, phycoerythrin, allophycocyanin, 6-carboxyfluorescein (6-FAM), 2',7'-dimethoxy-4',5'-dichloro-6-carboxyfluorescein (JOE), 6-carboxy-X-6-carboxy-2',4',7',4,7-hexachlorofluorescein (HEX), rhodamine(ROX), carboxyfluorescein (5-FAM) or N,N,N',N'-tetramethyl-6-carboxyrhodamine (TAMRA), radioactive labels, e.g. ³²P, ³⁵S, ³H; etc. The label may also be a two stage system, where the protein or (poly)peptide is conjugated to biotin, haptens, etc. having a high affinity binding partner, e.g. avidin, specific antibodies, etc., where the binding partner is conjugated to a detectable label. In another preferred embodiment of the present invention the label is a toxin, radioisotope, or fluorescent label.

In another preferred embodiment of the present invention, the (poly)peptide contains or lacks an N-terminal methionine. It is well known in the art that the N-terminal methionine encoded by the translation initiation codon generally is removed with high efficiency from any protein after translation in all eukaryotic cells. While the N-terminal methionine on most proteins also is efficiently removed in most prokaryotes, for some proteins this prokaryotic removal process is inefficient, depending on the nature of the amino acid to which the N-terminal methionine is covalently linked.

The present invention also relates to a protein complex comprising at least two proteins, wherein said at least two proteins are selected from the group of interaction partners listed in table 9. The term "protein complex" refers to a compound stably comprising at least two proteins. Preferably, said stability allows to purify said protein complex. In a preferred embodiment of the present invention, the protein complex comprises GIT1 and huntingtin.

The present invention also relates to the protein network of huntingtin, preferably the physical protein entities forming this network, which is described herein. In one embodiment, said protein network is formed by the interaction partners shown in table 6. Preferable, the protein network of the present invention is a validated protein network as described herein.

The present invention also relates to an antibody specifically recognizing the (poly)peptide of the present invention or specifically reacting with the protein complex of the present invention. This antibody is characterized in not recognizing the individual components of the protein complex but rather the complex itself. As such, said antibody recognizes a combined epitope, composed of amino acids of two different proteins within the protein complex. Dissociation of the complex will be detrimental to antibody recognition. Therefore, antibody binding depends on the integrity of the protein complex. In a preferred embodiment of the present invention, the antibody is specific for a protein complex comprising GIT1 and huntingtin.

In a preferred embodiment, the antibody of the present invention is polyclonal, monoclonal, chimeric, single chain, single chain Fv, human antibody, humanized antibody, or Fab fragment

In a more preferred embodiment of the present invention the antibody is labeled. Preferably, the label is selected from the group consisting of fluorochromes, e.g. fluorescein isothiocyanate (FITC), rhodamine, Texas Red, phycoerythrin, allophycocyanin, 6-carboxyfluorescein (6-FAM), 2',7'-dimethoxy-4',5'-dichloro-6-carboxyfluorescein (JOE), 6-carboxy-X-rhodamine(ROX), 6-carboxy-2',4',7',4,7-hexachlorofluorescein (HEX), 5-carboxyfluorescein (5-FAM) or N,N,N',N'-tetramethyl-6-carboxyrhodamine (TAMRA), radioactive labels, e.g. ³²P, ³⁵S, ³H; etc. The label may also be a two stage system, where the antibody is conjugated to biotin, haptens, etc. having a high affinity binding partner, e.g. avidin, specific antibodies, etc., where the binding partner is conjugated to a detectable label. In another preferred embodiment of the present invention the label is a toxin, radioisotope, or fluorescent label.

In a preferred embodiment of the present invention, the antibody is immobilized to a solid support. Preferably, the solid support may be the surface of a cell, a microtiter plate, beads or the surface of a sensor capable of detecting binding of the antibody or to the antibody.

The present invention also relates to a method of identifying whether a protein promotes huntingtin aggregation, comprising (a) transfecting a first cell with a nucleic acid molecule encoding a variant of the huntingtin protein or a fragment thereof capable of forming huntingtin aggregates; (b) co-transfecting a second cell with (i) a nucleic acid molecule encoding a variant of the huntingtin protein or a fragment thereof capable of forming huntingtin aggregates; and (ii) a nucleic acid molecule encoding a candidate modulator protein identified by the methods of the present invention or a nucleic acid molecule encoding a modulator protein selected from table 6 or table 7 (c) expressing the proteins encoded by the transfected nucleic acid molecule of (a) and (b); (d) isolating insoluble aggregates of huntingtin from the transfected cell of (a) and (b); and (e) determining the amount of insoluble huntingtin aggregates from the transfected cell of (a) and (b), wherein an increased amount of huntingtin aggregates isolated from the transfected cells of (b) in comparison with the amount of huntingtin aggregates isolated from the transfected cells of (a) is indicative of a protein's activity as an enhancer of huntingtin aggregation. Preferably, the huntingtin protein or protein fragment of step (a) is HD169Q68 or HD510Q68.

The present invention also relates to a method of identifying whether a protein inhibits huntingtin aggregation, comprising (a) transfecting a first cell with a nucleic acid molecule encoding a variant of the huntingtin protein or a fragment thereof capable of forming huntingtin aggregates; (b) co-transfecting a second cell with (i) a nucleic acid molecule encoding a variant of the huntingtin protein or a fragment thereof capable of forming huntingtin aggregates; and (ii) a nucleic acid molecule encoding a candidate modulator protein identified by the methods of the present invention or a nucleic acid molecule encoding a modulator protein selected from table 6 or table 7 (c) expressing the proteins encoded by the transfected nucleic acid molecule of (a) and (b); (d) isolating insoluble aggregates of huntingtin from the transfected cell of (a) and (b); and (e) determining the amount of insoluble huntingtin aggregates from the transfected cell of (a) and (b), wherein a reduced amount of huntingtin aggregates isolated from the transfected cells of (b) in comparison with the amount of huntingtin aggregates isolated from the transfected cells of (a) is indicative of a protein's activity as an inhibitor of huntingtin aggregation. Preferably, the huntingtin protein or protein fragment of step (a) is HD169Q68 or HD510Q68 or HdexQ51.

The term "promotes" means increasing the amount of huntingtin aggregation.

Preferably said huntingtin protein or the fragments thereof is selected from the proteins listed in table 6 and/or 7. Preferably said insoluble aggregates are isolated by using a filter retardation method comprising lysing cells and boiling in 2%SDS for 5min in the presence of 100mM DDT followed by a filtration step. The presence of aggregates is detected by using specific antibodies.

In a preferred embodiment of the present invention, determining the amount of insoluble huntingtin is performed by using light scattering or size exclusion chromatography. In another preferred embodiment of the present invention prior to step (d) the cells are treated with an ionic detergent. In yet another preferred embodiment of the methods of the present invention, the huntingtin aggregates are filtered or transferred onto a membrane.

The present invention also relates to a method for identifying compounds affecting, e.g. interfering or enhancing the interaction of huntingtin or of a direct or indirect

interaction partner of huntingtin comprising (a) contacting interacting proteins selected from the group of interacting proteins listed in table 6 in the presence or absence of a potential modulator of interaction; and (b) identifying compounds capable of modulating said interaction. The contacting is performed under conditions that permit the interaction of the two proteins. Sometimes more than two interacting proteins might be present in a single reaction as additional interaction partners of those listed under table 6, can be tested. However, the compound may also be a small molecule. Preferably said compounds are antibodies directed to huntingtin or to said interaction partner listed in table 6, wherein these antibodies are capable of interfering with the interaction with huntingtin. Alternatively, said compound is a peptide fragment of 10 to 25 amino acid residues of an interaction partner listed in table 7, wherein said peptide fragment is capable of interfering with the interaction with huntingtin. In a more preferred embodiment of the present invention, said antibody is an antibody directed to GIT1. In another more preferred embodiment of the invention, said peptide fragment is a peptide fragment of GIT1 of 10 to 25 capable of interfering with the interaction of GIT1 with huntingtin. Said interfering peptide may contain additional modifications in order to increase cellular uptake, solubility or to increase stability. Such modifications are known to the person skilled in the art and need not be listed here in detail. In a preferred embodiment of the present invention, the methods for identifying a compound further comprise the steps of modeling said compound by peptidomentics and chemically synthesizing the modeled compound.

In another preferred embodiment of the present invention, the methods for identifying a compound further comprise producing said compound. In yet another preferred embodiment of the present invention, the method for identifying said compound further comprise modifiying to achieve (i) modified site of action, spectrum of activity, organ specificity, and/or (ii) improved potency, and/or (iii) decreased toxicity (improved therapeutic index), and/or (iv) decreased side effects, and/or (v)

modified onset of therapeutic action, duration of effect, and/or (vi) modified pharmakinetic parameters (resorption, distribution, metabolism and excretion), and/or (vii) modified physico-chemical parameters (solubility, hygroscopicity, color, taste, odor, stability, state), and/or (viii) improved general specificity, organ/tissue specificity, and/or (ix) optimized application form and route by (i) esterification of

carboxyl groups, or (ii) esterification of hydroxyl groups with carbon acids, or (iii) esterification of hydroxyl groups to, e.g. phosphates, pyrophosphates or sulfates or hemi succinates, or (iv) formation of pharmaceutically acceptable salts, or (v) formation of pharmaceutically acceptable complexes, or (vi) synthesis of pharmacologically active polymers, or (vii) introduction of hydrophilic moieties, or (viii) introduction/exchange of substituents on aromates or side chains, change of substituent pattern, or (ix) modification by introduction of isosteric or bioisosteric moieties, or (x) synthesis of homologous compounds, or (xi) introduction of branched side chains, or (xii) conversion of alkyl substituents to cyclic analogues, or (xiii) derivatisation of hydroxyl group to ketales, acetales, or (xiv) N-acetylation to amides, phenylcarbamates, or (xv) synthesis of Mannich bases, imines, or transformation of ketones or aldehydes to Schiff's bases, oximes, acetales, ketales, enolesters, oxazolidines, thiozolidines or combinations thereof.

The present invention also relates to a method of diagnosing Huntington's disease in a biological sample comprising the steps of (a) contacting the sample with an antibody specific for a protein of table 6 or 7 or an antibody specific for the protein complex of the present invention; and (b) detecting binding of the antibody to a protein complex, wherein the detection of binding is indicative of Huntington's disease or of a predisposition to develop Huntington's disease. Preferably, binding is detected by measuring the presence of a fluorescent label bound to the protein complex.

In a preferred embodiment of the present invention's method protein complex contains (a) GIT1 or (b) said antibody is specific for a protein complex containing GIT1.

In a preferred embodiment of the present invention, said protein complex contains (a) at least one protein selected from htt, HIP15 or HP28 or (b) said antibody is specific for a protein complex containing at least one protein selected from htt, HIP15 or HP28.

The present invention also relates to a diagnostic agent/composition comprising the nucleic acid molecule of the present invention, the (poly)peptide of the present invention including/or the (poly)peptide mentioned in table 6 or 7, the antibody of the

present invention, an antibody specifically reacting with a protein selected from table 7 and/or a protein selected from table 7.

Moreover, the present invention also relates to a pharmaceutical composition comprising the nucleic acid molecule of the present invention, the (poly)peptide of the present invention, the interfering compound identified with a method of the present invention, the antibody of the present invention, an antibody specifically reacting with a protein selected from table 7 and/or a protein selected from table 7.

The pharmaceutical composition will be formulated and dosed in a fashion consistent with good medical practice, taking into account the clinical condition of the individual patient, the site of delivery of the pharmaceutical composition, the method of administration, the scheduling of administration, and other factors known to practitioners. The "effective amount" of the pharmaceutical composition for purposes herein is thus determined by such considerations.

As a general proposition, the total pharmaceutically effective amount of pharmaceutical composition administered parenterally per dose will be in the range of about 1 µg protein /kg/day to 10 mg protein /kg/day of patient body weight, although, as noted above, this will be subject to therapeutic discretion. More preferably, this dose is at least 0.01 mg protein /kg/day, and most preferably for humans between about 0.01 and 1 mg protein /kg/day for the peptide. If given continuously, the pharmaceutical composition is typically administered at a dose rate of about 1 µg/kg/hour to about 50 µg/kg/hour, either by 1-4 injections per day or by continuous subcutaneous infusions, for example, using a mini-pump. An intravenous bag solution may also be employed. The length of treatment needed to observe changes and the interval following treatment for responses to occur appears to vary depending on the desired effect.

Pharmaceutical compositions of the invention may be administered orally, rectally, parenterally, intracistemally, intravaginally, intraperitoneally, topically (as by powders, ointments, drops or transdermal patch), bucally, or as an oral or nasal spray. By "pharmaceutically acceptable carrier" is meant a non-toxic solid, semisolid or liquid filler, diluent, encapsulating material or formulation auxiliary of any type. The term "parenteral" as used herein refers to modes of administration which include

intravenous, intramuscular, intraperitoneal, intrasternal, subcutaneous and intraarticular injection and infusion.

The pharmaceutical composition is also suitably administered by sustained-release systems. Suitable examples of sustained-release compositions include semipermeable polymer matrices in the form of shaped articles, e.g., films, or Sustained-release matrices include polylactides (U.S. Pat. No. mirocapsules. 3,773,919, EP 58,481), copolymers of L-glutamic acid and gamma-ethyl-L-glutamate (Sidman, U. et al., Biopolymers 22:547-556 (1983)), poly (2- hydroxyethyl methacrylate) (R. Langer et al., J. Biomed. Mater. Res. 15:167-277 (1981), and R. Langer, Chem. Tech. 12:98-105 (1982)), ethylene vinyl acetate (R. Langer et al., Id.) or poly-D- (-)-3-hydroxybutyric acid (EP 133,988). Sustained-release pharmaceutical composition also include liposomally entrapped protein, antibody, (poly)peptide, peptide or nucleic acid. Liposomes containing the pharmaceutical composition are prepared by methods known per se: DE 3,218,121; Epstein et al., Proc. Natl. Acad. Sci. (USA) 82:3688-3692 (1985); Hwang et al., Proc. Natl. Acad. Sci. (USA) 77:4030-4034 (1980); EP 52,322; EP 36,676; EP 88,046; EP 143,949; EP 142,641; Japanese Pat. Appl. 83-118008; U.S. Pat. Nos. 4,485,045 and 4,544,545; and EP 102,324. Ordinarily, the liposomes are of the small (about 200-800 Angstroms) unilamellar type in which the lipid content is greater than about 30 mol. percent cholesterol, the selected proportion being adjusted for the optimal therapy.

For parenteral administration, in one embodiment, the pharmaceutical composition is formulated generally by mixing it at the desired degree of purity, in a unit dosage injectable form (solution, suspension, or emulsion), with a pharmaceutically acceptable carrier, i.e., one that is non-toxic to recipients at the dosages and concentrations employed and is compatible with other ingredients of the formulation. For example, the formulation preferably does not include oxidizing agents and other compounds that are known to be deleterious to (poly)peptides.

Generally, the formulations are prepared by contacting the components of the pharmaceutical composition uniformly and intimately with liquid carriers or finely divided solid carriers or both. Then, if necessary, the product is shaped into the desired formulation. Preferably the carrier is a parenteral carrier, more preferably a solution that is isotonic with the blood of the recipient. Examples of such carrier

vehicles include water, saline, Ringer's solution, and dextrose solution. Non-aqueous vehicles such as fixed oils and ethyl oleate are also useful herein, as well as liposomes. The carrier suitably contains minor amounts of additives such as substances that enhance isotonicity and chemical stability. Such materials are nontoxic to recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate, succinate, acetic acid, and other organic acids or their salts; antioxidants such as ascorbic acid; low molecular weight (less than about ten residues) (poly)peptides, e.g., polyarginine or tripeptides; proteins, such as serum hydrophilic immunoglobulins; polymers such gelatin, albumin, or polyvinylpyrrolidone; amino acids, such as glycine, glutamic acid, aspartic acid, or arginine; monosaccharides, disaccharides, and other carbohydrates including cellulose or its derivatives, glucose, manose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sorbitol; counterions such as sodium; and/or nonionic surfactants such as polysorbates, poloxamers, or PEG. The proteinacous components of the pharmaceutical composition are typically formulated in such vehicles at a concentration of about 0.1 mg/ml to 100 mg/ml, preferably 1-10 mg/ml, at a pH of about 3 to 8. It will be understood that the use of certain of the foregoing excipients, carriers, or stabilizers will result in the formation protein or (poly)peptide salts.

The components of the pharmaceutical composition to be used for therapeutic administration must be sterile. Sterility is readily accomplished by filtration through sterile filtration membranes (e.g., 0.2 micron membranes). Therapeutic components of the pharmaceutical composition (poly)peptide compositions generally are placed into a container having a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle.

The components of the pharmaceutical composition ordinarily will be stored in unit or multi-dose containers, for example, sealed ampoules or vials, as an aqueous solution or as a lyophilized formulation for reconstitution. As an example of a lyophilized formulation, 10-ml vials are filled with 5 ml of sterile-filtered 1% (w/v) aqueous protein solution, and the resulting mixture is lyophilized. The infusion solution is prepared by reconstituting the lyophilized protein using bacteriostatic Water-for-Injection.

The invention also provides a pharmaceutical/diagnostic pack or kit comprising one or more containers filled with one or more of the ingredients of the pharmaceutical/diagnostic compositions of the invention. Associated with such container(s) can be a notice in the form prescribed by a governmental agency regulating the manufacture, use or sale of pharmaceuticals or biological products, which notice reflects approval by the agency of manufacture, use or sale for human administration. In addition, the (poly)peptides of the components of the pharmaceutical composition invention may be employed in conjunction with other therapeutic compounds.

Finally, the present invention relates to the use of the nucleic acid molecule of the present invention, the interfering compound identified with a method of the present invention, the (poly)peptide of the present invention including/or the (poly)peptide mentioned in table 6 or 7, the antibody of the present invention, an antibody specifically reacting with a protein selected from table 7 and/or a protein selected from table 7 for the preparation of a pharmaceutical composition for the treatment of Huntington's disease.

Tables:

Table 1:

PROTEIN-PROTEIN INTERACTIONS IN	
THE PPI OF HUNTINGTIN	
Baits (DBD)	Preys (AD)
BARD1	PLIP
EF1G	EF1G
HD1.7	CA150
HD1.7	HIP1
HD1.7	HYPA
HD1.7	SH3GL3
HDexQ20	CA150
HDexQ20	НҮРА
HDexQ20	SH3GL3
HDexQ51	CA150
HDexQ51	НҮРА
HDexQ51	SH3GL3
mp53	p53
mp53	PIASy
PIASy	SUMO-2
PIASy	SUMO-3
VIM	NEFL
VIM	VIMc
BARD1	BAIP1
BARD1 ·	BAIP2
BARD1	BAIP3
BARD1	FEZ1
BARD1	GIT1
BARD1	HBO1
BARD1	HIP5
BARD1	HZFH
BARD1	IKAP
BARD1	mHAP1
BARD1	NAG4
BARD1	PIASy
BARD1	PTN ·
BARD1	SETBD1

BARD1	ZHX1
CLH-17	Ku70
CLH-17	PIASy
GADD45G	BAIP3
GADD45G	CGI-125
GADD45G	CGI-723
	EF1A
GADD45G	
GADD45G	EF1G
GADD45G	G45IP1
GADD45G	G45IP2
GADD45G	G45IP3
GADD45G	HIP16
GADD45G	HIP5
GADD45G	LUC7B1
GADD45G	PIASy
GADD45G	PLIP
GADD45G	PTN
GADD45G	PTPK
hADA3	BAIP1
hADA3	Ku70
hADA3	MAGEH1
hADA3	PIASy
HD1.7	CGI-125
HD1.7	DRP-1
HD1.7	FEZ1
HD1.7	GIT1
HD1.7	HIP11
HD1.7	HIP13
HD1.7	HIP15
HD1.7	HIP16
HD1.7	HIP5
HD1.7	HZFH
HD1.7	IKAP
HD1.7	Ku70
HD1.7	PIASy
HDd1.0	FEZ1 .
HDd1.0	GIT1
HDd1.0	IKAP
HDd1.3	HZFH

HDd1.3	IKAP
HDd1.3	Ku70
HDd1.3	PIASy
HDexQ20	CGI-125
HDexQ20	HIP13
HDexQ20	HP28
HDexQ20	PFN2
HDexQ51	CGI-125
HDexQ51	HIP13 .
HDexQ51	HIP15
HDexQ51	HP28
HDexQ51	PFN2
HIP2	PIASy
HIP5	APP1
	BAIP1
HIP5	BAIP2
HIP5	CGI-74
	FEZ1
HIP5	GIT1
HIP5	
HIP5	HBO1
HIP5	HMP
HIP5	KPNA2
HIP5	mHAP1
HIP5	NAG4
HIP5	PLIP
IMPD2	PIASy.
KPNB1	PIASy
KPNB1	PTN
mp53	HZFH
mp53	ZHX1
PIASy	MAPIc3
TAL1	ZHX1
TCP1G	Ku70
VIM	ALEX2
VIM	BAIP1
VIM	DRP-1
VIM	G45IP1
VIM ·	HBO1
VIM .	HSPC232

VIM	HZFH
VIM	PIASy
VIM	SETBD1
VIM .	SH3GL3
ZNF33B	mHAP1
ZNF33B	ZHX1 ·

Table 2 Classification of proteins in Huntington's disease interaction network FUSION ACCESSION IDEN aa MATCH LOC ID NAME Huntingtin fragments HD1.7 1-506 N, C DBD 100 P42858 huntingtin N, C DBD P42858 100 1-320 HDd1.0 huntingtin DBD P42858 100 166-506 N, C HDd1.3 huntingtin 1-90 N, C HdexQ20 huntingtin DBD P42858 96 DBD P42858 75 1-82 N, C HdexQ51 huntingtin Transcriptional control and DNA maintenance DBD Q99728 99 1-379 N BRCA1 associated ring domain protein 1 BARD1 93 299-629 N AD 014776 CA150 putative transcription factor CA150 N DBD 095257 100 18-159 GADD45G growth arrest and DNA damage inducible protein GADD45 gamma Ν DBD 075528 100 235-432 hADA3 ADA3 like protein 100 1-611 Ν HBO1 histone acetyltransferase binding to ORC AD 095251 5-510 N, C AD, DBD Q8N2W9 100 **PIASy** protein inhibitor of activated STAT protein gamma (PIASy) 075400 100 8-422 C, N huntingtin interacting protein HYPA/FBP11 (fragment) ΑD **HYPA** 100 1830-2000 Ν AD, DBD Q9Y4I0 **HZFH** zinc finger helicase HZFH N, C 1207-1332 AD 095163 100 IKAP IKK complex associated protein 298-608 Ν AD P12956 100 ATP dependent DNA helicase II, 70 kDa subunit Ku70 94-651 bromodomain containing protein NAG4 AD Q9NPI1 100 N NAG4 AD P04637 100 1-393 N p53 cellular tumor antigen p53 100 248-393 Ν cellular tumor antigen p53 (C-terminus) AD P04637 p53c cellular tumor antigen p53 (mouse) DBD P02340 100 73-390 N mp53 ΑD O95624 100 5-461 N, PN PLIP cPLA? interacting protein AD Q15047 100 1023-1291 Ν SETDB1 histone-lysine N-methyltransferase, H3 lysine-9 specific 4 P55854 100 1-103 C, N ΑD ubiquitin like protein SMT3A (SUMO-2) SUMO-2 C, N 1-95 SUMO-3 ubiquitin like protein SMT3B (SUMO-3) ΑD P55855 100 145-873 Ν zinc finger homeobox protein ZHX1 AD Q9UKY1 100 ZHX1 527-778 Ν DBD Q8NDW3 100 zinc finger protein 33b ZNF33B Cellular organization and protein transport P51693 100 243-555 PM, EC AD APP1 amyloid like protein 1 precursor 1-289 PM, V Q00610 100 dathrin heavy chain 1 DBD CLH-17 axonemal dynein light chain (hp28) AD Q9BQZ6 100 3-258 CN HP28 O35668 100 3-471 C, EE ΑD mHAP1 huntingtin associated protein 1 (mouse) C, GN HIP1 AD 000291 100 245-631 huntingtin interacting protein 1 Q16891 100 212-758 Mit mitofilin AD **HMP** microtubule associated proteins 1A/1B light chaln 3 ΑD Q9H491 100 58-170 CN, MT MAP1Ic3 Q8IU72 100 1-543 CN, IF light molecular weight neurofilament protein AD **NEFL**

SH3GL3 SH3 containing GRB2 like protein 3 AD Q99963 100 3-347 KPNA2 karyopherin alpha-2 subunit AD P52292 100 141-529	CN M, EC V C, N C, N CN, IF
SH3GL3 SH3 containing GRB2 like protein 3 AD Q99963 100 3-347 KPNA2 karyopherin alpha-2 subunit AD P52292 100 141-529	V C, N C, N CN, IF
KPNA2 karyopherin alpha-2 subunit AD P52292 100 141-529	C, N C, N CN, IF
Turing the population of the p	C, N CN, IF
KPNR1 karvopherin beta-1 subunit DBD Q14974 100 668-876	N, IF
18 Ltm. Designation of London	-
VIM vimentin DBD P08670 100 1-466 0	N, IF
VIMc vimentin (C-terminus) AD P08670 100 190-466 C	
Cell signaling and fate	
	C, PM
CLK1 protein kinase CLK1 DBD P49759 100 209-484	N
FEZ1 fasciculation and elongation protein zeta 1 AD Q99689 100 131-392 0	C, PM
GIT1 ARF GTPese activating protein GIT1 AD Q9Y2X7 98 249-761 I	PM, V
PTPK protein-tyrosine phosphatase kappa precursor AD Q15262 100 1227-1439 F	M, AJ
Cellular metabolism	
DRP-1 dihydropyrimidinase related protein 1 (C-terminus) AD Q14194 100 345-572	С
IMPD2 inosine-5'-monophosphate dehydrogenase 2 DBD P12268 100 34-514	С
TAL1 transaldolase DBD P37837 100 3-337	<u> </u>
Protein synthesis and turnover	
EF1A translation elongation factor 1 alpha 1 · AD P04720 100 294-462	C, MT
EF1G elongation factor 1 gamma AD, DBD P26641 100 2-437	C, MT
EF1Gc elongation factor 1 gamma (C-terminus) AD P26641 100 123-437	C, MT
HIP2 ubiquitin conjugating enzyme E2-25 kDa DBD P27924 100 1-200	C, N
TCPG T-complex protein 1, gamma subunit DBD P49368 100 252-544	<u>C</u>
Uncharacterized proteins	
BAIP1 BARD1 interacting protein 1[similar to RIKEN cDNA 1810018M11] AD Q9BS30 100 1-226	UN
BAIP2 BARD1 interacting protein 2 [hypothetical protein] AD Q9H0I6 100 107-684	UN
BAIP3 BARD1 interacting protein 3 [hypothetical protein] AD Q96HT4 100 152-436	UN
CGI-74 CGI-74 protein AD Q9Y383 100 159-270	·UN
CGI-125 CGI-125 protein AD Q9Y3C7 100 1-131	UN
G45iP1 GADD45G interacting protein 1[hypothetical protein] AD Q9H0V7 100 1-340	UN
G45IP2 GADD45G interacting protein 2 [B2 gene partial cDNA, clone B2E] AD Q9NYA0 100 566-926	UN
G45iP3 GADD45G interacting protein 3 [OK/SW-CL.16] AD Q8Ni70 100 3-134	ÜN
HIP5 huntingtin interacting protein 5 [hypothetical protein KIAA1377] AD, DBD Q9P2H0 100 445-988	N, C
HIP11 huntingfin interacting protein 11[hypothetical protein] AD Q96EZ9 100 176-328	UN
HIP13 huntingtin interacting protein 13 [metastasis suppressor protein] AD Q96RX2 100 512-755	UN
HIP15 huntingtin interacting protein 15 [similar to KIAA0443 gene product] AD Q96D09 100 663-638	UN
HIP16 huntingtin interacting protein 16 [similar to KIAA0266 gene product] AD Q9BVJ6 100 585-771	UN
HSPC232 HSPC232 AD Q9P0P6 92 1-319	UN
LUC7B1 putative SR protein LUC7B1 (SR+89) AD Q9NQ29 99 116-371	ER
MAGEH1 melanoma associated antigen H1 AD Q9H213 100 1-219	UN

Abbreviations: aa, amino acids; IDEN, Identity; LOC, localisation; AD, activation domain; DBD, DNA binding domain; AJ, adherens junctions; C, cytosol; CN, cytoskeleton; EC, extracellular space; EE, early endosomes; ER, endoplasmic reticulum; IF, intermediate filaments; GN, Golgi network; Mit, mitochondria; MT, microtubules; N, nucleus; PM, plasma membrane; PN, perinuclear; UN, unknown; V, vesicles; [], database annotation

Table 3 New proteins in Huntington's disease interaction network

ID	NAME	FUSION	ACCESSION	IDEN	aa MATCH	LOC
Transcript	ional control and DNA maintenance					
BARD1	BRCA1 associated ring domain protein 1	DBD	Q99728	99	1-379	N
CA150	putative transcription factor CA150	AD	O14776	93	299-629	N
Cell signa	ling and fate					
GIT1	ARF GTPase activating protein GIT1	AD	Q9Y2X7	98	249-761	PM, V
HSPC232	HSPC232	AD	Q9P0P6	92	1-319	UN
LUC7B1	putative SR protein LUC7B1 (SR+89)	AD	Q9NQ29	99	116-371	ER

Abbreviations: aa, amino acids; IDEN, identity; LOC, localisation; AD, activation domain; DBD, DNA binding domain; AJ, adherens junctions; C, cytosol; CN, cytoskeleton; EC, extracellular space; EE, early endosomes; ER, endoplasmic reticulum; IF, intermediate filaments; GN, Golgi network; Mit, mitochondria; MT, microtubules; N, nucleus; PM, plasma membrane; PN, perinuclear; UN, unknown; V, vesicles; [], database annotation

Table 4:

New protein-protein inter	actions, found
Baits (DBD)	Preys (AD)
BARD1	BAIP1
BARD1	BAIP2
BARD1	BAIP3
BARD1	FEZ1
BARD1	GIT1
BARD1	HBO1
BARD1	HIP5
BARD1	HZFH
BARD1	IKAP
BARD1	mHAP1
BARD1	NAG4
BARD1	PIASy ·
BARD1	PTN
BARD1	SETBD1
BARD1	ZHX1
CLH-17	Ku70
CLK1	PIASy
GADD45G	BAIP3
GADD45G	CGI-125
GADD45G	CGI-74

CADDAEC	EF1A
GADD45G	
GADD45G	EF1G
GADD45G	G45IP1
GADD45G	G45IP2
GADD45G	G45IP3
GADD45G	HIP16
GADD45G	HIP5
GADD45G	LUC7B1
GADD45G	PIASy
GADD45G	PLIP
GADD45G	PTN
GADD45G	PTPK
hADA3	BAIP1
hADA3	Ku70
hADA3	MAGEH1
hADA3	PIASy
HD1.7	CGI-125
HD1.7	DRP-1
HD1.7	FEZ1
HD1.7	GIT1
HD1.7	HIP11
HD1.7	HIP13
HD1.7	HIP15
HD1.7	HIP16
HD1.7	HIP5
HD1.7	HZFH
HD1.7	IKAP
HD1.7	Ku70
HD1.7	PIASy
HDd1.0	FEZ1
HDd1.0	GIT1
HDd1.0	IKAP .
HDd1.3	HZFH
HDd1.3	IKAP
HDd1.3	Ku70
HDd1.3	PIASy
HDexQ20	CGI-125
HDexQ20	HIP13
HDexQ20	HP28

HDexQ51	HDexQ20	PFN2
HDexQ51	 	
HDexQ51 HIP15 HDexQ51 HP28 HDexQ51 PFN2 HIP2 PFN2 HIP5 APP1 HIP5 BAIP1 HIP5 BAIP2 HIP5 BAIP2 HIP5 GIT4 HIP5 FEZ1 HIP5 HBO1 HIP5 HMP HIP5 HMP HIP5 MAC4 HIP5 MAC4 HIP5 NAG4 HIP5 NAG4 HIP5 PLIP IMP02 PIASY KPNB1 PTN mp53 HZFH mp53 HZFH mp53 ZHX1 PIASY MAPIc3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM HBO1 VIM HBO1 VIM HSPC232		
HDexQ51 HP28 HDexQ51 PFN2 HIP2 PIASy HIP5 APP1 HIP5 BAIP1 HIP5 BAIP2 HIP5 BAIP2 HIP5 CGI-74 HIP5 FEZ1 HIP5 HBO1 HIP5 HBO1 HIP5 HMP HIP5 MAP1 HIP5 MAP4 HIP5 NAG4 HIP5 NAG4 HIP5 PLIP IMPD2 PIASY KPNB1 PTN mp53 HZFH mp53 HZFH mp53 ZHX1 PIASY MAPIC3 TAL1 ZHX1 TCP1G Ku70 VIM BAIP1 VIM DRP-1 VIM BAIP1 VIM HBO1 VIM HBO1 VIM HSPC232 VIM HSPC33B		
HDexQ51 HIP2 HIP5 HIP5 APP1 HIP5 BAIP1 HIP5 BAIP2 HIP5 BAIP2 HIP5 CGI-74 HIP5 FEZ1 HIP5 HIP5 HB01 HIP5 HB01 HIP5 HMP HIP5 HIP5 NAG4 HIP5 NAG4 HIP5 PLIP IMPD2 PIASY KPNB1 PTN mp53 PTN mp53 TAL1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM VIM CG45IP1 VIM VIM VIM PIASY VIM VIM PIASY VIM VIM PIASY PI		
HIP2		
HIP5		
HIP5 BAIP1 HIP5 BAIP2 HIP5 CGI-74 HIP5 FEZ1 HIP5 GIT1 HIP5 HBO1 HIP5 HMP HIP5 KPNA2 HIP5 MAG4 HIP5 NAG4 HIP5 PLIP IMPD2 PIASY KPNB1 PIASY KPNB1 PTN mp53 TAL1 ZHX1 TCP1G Ku70 VIM BAIP1 VIM G45IP1 VIM G45IP1 VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1		
HIP5		BAIP1
HIP5 FEZ1 HIP5 FEZ1 HIP5 GIT1 HIP5 HB01 HIP5 HMP HIP5 KPNA2 HIP5 MHAP1 HIP5 NAG4 HIP5 PLIP IMPD2 PIASY KPNB1 PIN mp53 HZFH mp53 ZHX1 PIASY MAPIC3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM G45IP1 VIM HB01 VIM HSPC232 VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B MHAP1		
HIP5 FEZ1 HIP5 GIT1 HIP5 HBO1 HIP5 HMP HIP5 HMP HIP5 KPNA2 HIP5 MHAP1 HIP5 NAG4 HIP5 PLIP IMPD2 PIASY KPNB1 PIN mp53 HZFH mp53 ZHX1 PIASY MAPIC3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM G45IP1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM PIASY VIM PIASY VIM PIASY VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B MHAP1	HIP5	CGI-74
HIP5 HMP HIP5 HMP HIP5 KPNA2 HIP5 MHAP1 HIP5 MAG4 HIP5 PLIP IMPD2 PIASY KPNB1 PIN mp53 HZFH mp53 ZHX1 PIASY MAPIc3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM G45IP1 VIM G45IP1 VIM HSPC232 VIM HZFH VIM PIASY VIM PIASY VIM PIASY VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B MHAP1	HIP5	FEZ1
HIP5	HIP5	GIT1
HIP5	HIP5	HBO1
HIP5	HIP5	HMP
HIP5	HIP5	KPNA2
HIP5 PLIP IMPD2 PIASy KPNB1 PIASy KPNB1 PTN mp53 HZFH mp53 ZHX1 PIASy MAPIc3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HZFH VIM PIASy VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	HIP5	mHAP1
IMPD2 PIASy KPNB1 PIASy KPNB1 PTN mp53 HZFH mp53 ZHX1 PIASY MAPIc3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	HIP5	NAG4
IMPD2 PIASY KPNB1 PIASY KPNB1 PTN mp53 HZFH mp53 ZHX1 PIASY MAPIc3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	HIP5	PLIP
KPNB1 PTN mp53 HZFH mp53 ZHX1 PIASy MAPIc3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASy VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	IMPD2	PIASy .
mp53 HZFH mp53 ZHX1 PIASy MAPIc3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	KPNB1	PIASy
mp53 ZHX1 PIASy MAPIc3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASy VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	KPNB1	PTN
PIASY MAPIc3 TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASy VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	mp53	HZFH
TAL1 ZHX1 TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	mp53	ZHX1
TCP1G Ku70 VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASy VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	PIASy	MAPIc3
VIM ALEX2 VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	TAL1	ZHX1
VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASy VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	TCP1G	Ku70
VIM DRP-1 VIM G45IP1 VIM HBO1 VIM HSPC232 VIM HZFH VIM PIASy VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	VIM	ALEX2
VIM G45IP1 VIM HB01 VIM HSPC232 VIM HZFH VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	VIM	BAIP1
VIM HB01 VIM HSPC232 VIM HZFH VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	VIM	DRP-1
VIM HSPC232 VIM HZFH VIM PIASy VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	VIM	G45IP1
VIM HZFH VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	VIM .	HBO1
VIM PIASY VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	VIM	HSPC232
VIM SETBD1 VIM SH3GL3 ZNF33B mHAP1	VIM	HZFH
VIM SH3GL3 ZNF33B mHAP1	VIM	PIASy
ZNF33B mHAP1	VIM	SETBD1
	VIM	SH3GL3
7NE22D 7UV4	ZNF33B	mHAP1
ZINFOOD ZMXI	ZNF33B	ZHX1

Table 5:

- * Aarskog syndrome
- * Achromatopsia
- * Acoustic neuroma
- * Adrenal hyperplasia
- * Adrenoleukodystrophy
- * Agenesis of corpus callosum
- * Aicardi syndrome
- * Alagille syndrome
- * Albinism
- * Alopecia areata
- * Alstrom syndrome
- * Alpha-1-antitrypsin deficiency
- * Alzheimer
- * Ambiguous genitalia
- * Androgen insensitivity syndrome(s)
- * Anorchia
- * Angelman syndrome
- * Anopthalmia
- * Apert syndrome
- * Arthrogryposis
- * Ataxia
- * Autism
- * Bardet-Biedl syndrome
- * Basal cell carcinoma
- * Batten disease
- * Beckwith-Wiedemann syndrome
- * Blepharophimosis
- * Blind
- * Branchio-Oto-Renal (BOR) syndrome
- * Canavan
- * Cancer: (ataxia telangiectasia, basal cell nevus, brain /spine, breast, colon / bowel, leukemia / lymphoma, lung, melanoma / skin, multiple endocrine neoplasia, oral, ovarian, prostate, retinoblastoma, testicular, von Hippel-Lindau, xeroderma pigmentosa)
- * Cardiof ciocutaneous syndrome
- * Celiac sprue
- * Charcot-Marie-Tooth
- * CHARGE association
- * Chromosome anomalies trisomy, deletions, inversions, duplications, translocations, 4p- (Wolf-Hirshhorn), 5 (cri-du-chat, 5p-), 6, 8p, 9 (trisomy 9, 9p-), 11 (11q, 11;22), 13 (trisomy 13, Patau), 15, 16 (mosaic), 18 (18q-, 18p-, ring 18, trisomy 18, tetrasomy 18p, Edwards), 21 (Down syndrome, trisomy 21), 22, X & Y [sex chromosome anomalies, Klinefelter (XXY, other), Turner (XO, other), fragile-X, other]
- * Cleft lip and/or cleft palate
- * Cockayne syndrome
- * Coffin-Lowry syndrome
- * Coffin-Siris syndrome
- * Congenital heart defects

- * Connective tissue conditions
- * Cooley anemia
- * Conjoined twins
- * Cornelia de Lange syndrome
- * Costello syndrome
- * Craniofacial conditions
- * Cri-du-Chat (5p-)
- * Cystic fibrosis
- * Cystinosis
- * Cystinuria
- * Dandy-Walker syndrome
- * Deaf / hard of hearing
- * Dermatological (skin) conditions
- * Developmental delay / mental retardation
- * DiGeorge syndrome
- * Down syndrome
- * DRPLA
- * Dubowitz syndrome
- * Dwarfism/ short stature
- * Dysautonomia
- * Dystonia
- * Ectodermal dysplasia
- * Ehlers Danlos syndrome
- * Endocrine Conditions
- * Epidermolysis bullosa
- * Facial anomalies, disfigurement
- * Fanconi anemia
- * Fetal alcohol syndrome and effects
- * FG syndrome
- * Fragile-X syndrome
- * Friedreich ataxia
- * Freeman Sheldon syndrome
- * Galactosemia
- * Gardner syndrome
- * Gastroenterology conditions
- * Gaucher disease
- * Glycogen storage disease
- * Goldenhar syndrome
- * Gorlin syndrome
- * Hallerman Streiff syndrome
- * Hearing problems
- * Heart conditions
- * Hemochromatosis
- * Hemophilia
- * Hemoglobinopathies
- * Hereditary hemorrhagic telangiectasia
- * Hereditary spastic paraplegia
- * Hermansky-Pudlak syndrome
- * Hirschsprung anomaly
- * Holoprosencephaly

- * Huntington disease
- * Hydrocephalus
- * Ichthyosis
- * Immune deficiencies
- * Incontinentia pigmenti
- * Infertility
- * Intestinal problems
- * Joseph disease
- * Joubert syndrome
- * Kabuki syndrome
- * Kidnev conditions
- * Klinefelter syndrome
- * Klippel-Feil syndrome
- * Klippel-Trenaunay syndrome
- * Langer-Giedion syndrome
- * Laurence-Moon-Biedl syndrome
- * Leber Optic Atrophy
- * Leigh disease
- * Lesch-Nyhan syndrome
- * Leukodystrophy [Adrenoleukodystrophy (ALD), Alexanders Disease, CADASIL (Cerebral Autosomal Dominant Arteriopathy with Subcortical Infarcts & Leukoencephalopathy), Canavan Disease (Spongy Degeneration), Cerebrotendinous Xanthomatosis (CTX), Globoid Cell (Krabbes) Leukodystrophy, Metachromatic Leukodystrophy (MLD), Ovarioleukodystrophy, Pelizaeus-Merzbacher Disease, Refsum Disease, van der Knaap syndrome, Zellweger syndrome]
- * Limb anomalies [missing arm(s) or leg(s), Poland anomaly, other]
- * Lissencephaly [Isolated Sequence (ILS), X-Linked (XLIS), Subcortical Band Heterotopia (SBH), Miller-Dieker syndrome (MDS), Microcephaly, Microlissencephaly (MLIS), Norman-Roberts syndrome (NRS), With Cerebellar Hypoplasia (LCH), Polymicrogyria (PMG), Schizencephaly (SCH), Muscle-Eye-Brain (MEB) Disease, and Walker-Warburg syndrome (WWS), 17p13.3 deletion]
- * Liver conditions (biliary atresia, Alagille syndrome, alpha-1 antitrypsin, tyrosinemia, neonatal hepatitis, Wilson disease)
- * Lowe syndrome
- * Lung / pulmonary conditions
- * Lymphedema
- * Maffucci syndrome(Ollier, multiple cartilaginous enchondromatosis)
- * Malignant hyperthermia
- * Maple syrup urine disease
- * Marinesco-Sjogren Syndrome
- * Marfan syndrome
- * Menke syndrome
- * Mental retardation / developmental delay
- * Metabolic conditions (carbohydrate deficient glycoprotein syndrome (CDGS), diabetes insipidus, Fabry, galactosemia, glucose-6-phosphate dehydrogenase (G6PD), fatty acid oxidation disorders, glutaric aciduria, hypophosphatemia, Krabbe, lactic acidosis, lysosomal storage diseases, mannosidosis, maple syrup urine, mitochondrial, neuro-metabolic, organic acidemias, PKU, purine, pyruvate dehydrogenase deficiency, urea cycle conditions, vitamin D deficient rickets)

- * Miscarriage, stillbirth, infant death
- * Mitochondrial conditions (Alpers, Barth, beta-oxidation defects, carnitine deficiency, CPEO, Keams-Sayre, lactic acidosis, Leber optic neuropathy, Leigh, LCAD, Luft, MCAD, MAD, glutaric aciduria, MERRF, MNGIE, NARP, Pearson, PHD, SCAD, NADH-CoQ reductase, succinate dehydrogenase, Complex III, Complex IV, COX, Complex V, other)
- * Moebius syndrome
- * Mucolipidosis, type IV (ML4)
- * Mucopolysaccharidosis (Hunter syndrome, Hurler syndrome, Maroteaux-Lamy syndrome, Sanfilippo syndrome, Scheie syndrome, Morquio syndrome, other)
- * Multiple hereditary exostoses
- * Muscular dystrophy /atrophy (neuromuscular conditions including: Duchenne, facioscapulohumeral, Charcot Marie Tooth, spinal muscular atrophy, other)
- * Myotonic dystrophy
- * Nager & Miller syndromes
- * Nail Patella syndrome
- * Narcolepsy
- * Neurologic conditions (neuro-metabolic, neurogenetics, neuromuscular, other)
- * Neurofibromatosis (von Recklinghausen)
- * Neuromuscular conditions
- * Niemann-Pick disease
- * Noonan syndrome
- * Opitz syndromes [Opitz-Frias, Opitz FG (Opitz-Kaveggia), Opitz-C (Trigonocephaly)]
- * Organic acidemias
- * Osler-Weber-Rendu syndrome
- * Osteogenesis imperfecta
- * Oxalosis & hyperoxaluria
- * Pallister-Hall syndrome
- * Pallister-Killian syndrome (tetrasomy 12p, Teschler-Nicola syndrome)
- * Parkinson's disease
- * Periodic paralysis
- * Phenylketonuria (PKU)
- * Polycystic kidney disease
- * Popliteal pterygium syndrome
- * Porphyria
- * Prader-Willi syndrome
- * Progeria (Werner, Hutchinson-Gilford, Cockayne, Rothmond-Thomson syndromes)
- * Proteus syndrome
- * Prune belly syndrome
- * Pseudoxanthoma elasticum (PXE)
- * Psychiatric conditions
- * Refsum disease
- * Retinal degeneration
- * Retinitis pigmentosa (retinal degenerative diseases, Usher syndrome)
- * Retinoblastoma
- * Rett syndrome
- * Robinow syndrome
- * Rubinstein-Taybi syndrome
- * Russell-Silver syndrome

- * SBMA
- * SCA
- * Schizencephaly
- * Sex chromosome anomalies (47,XXY, 47,XXX, 45,X and variants, 47,XYY)
- * Shwachman syndrome
- * Sickle cell anemia
- * Skeletal dysplasia
- * Smith-Lemli-Opitz syndrome (RHS syndrome)
- * Smith-Magenis syndrome (17p-)
- * Sotos syndrome
- * Spina bifida (myelomeningocele, neural tube defects)
- * Spinal muscular atrophy (Werdnig-Hoffman, Kugelberg-Welander)
- * Stickler / Marshall syndrome
- * Sturge-Weber
- * Tay-Sachs disease / other (dysautonomia, dystonia, Gaucher, Niemann Pick, Canavan, Bloom)
- * Thalassemia (Cooley anemia)
- * Thrombocytopenia absent radius syndrome
- * Tourette syndrome
- * Treacher Collins syndrome (craniofacial)
- * Trisomy (21, 18, 13, 9, other, see chromosome syndromes)
- * Tuberous sclerosis
- * Turner syndrome
- * Twins / triplets / multiple births
- * Unknown disorders
- * Urea cycle conditions
- * Usher syndrome
- * VATER association
- * Velo-cardio-facial syndrome (Shprintzen, DiGeorge, 22q deletion)
- * Visual impairment / blind
- * Von Hippel-Lindau syndrome
- * Waardenburg syndrome
- * Weaver syndrome
- * Werner syndrome
- * Williams syndrome
- * Wilson disease (hepatolenticular degeneration)
- * Xeroderma pigmentosum
- * Zellweger syndrome

TABLE 6

PROTEIN-PROTEIN INTERACTIONS IN THE PROTEIN NETWORK OF HUNTINGTIN

PROTEIN-PRO	TEIN INTER
BAIT	PREY
SETDB1	SUMO-3
PIASy	SUMO-3
HZFH	SUMO-3
PIASy	HYPA
HZFH	HYPA
MAP1lc3	HYPA
ZHX1	HYPA
PIASy	HZFH
HZFH	HZFH
GIT1	HZFH
VIM	HZFH
PIASy	ZHX1
HZFH	ZHX1
VIM	ZHX1
FEZ1	HMP
HZFH	HMP
НМР	HMP
PIASy	HMP
HZFH .	PTN
HIP15	PTN ·
PIASy	PTN
PTN	PTN
FEZ1	PTN
KPNA2	G45IP3
GIT1	G45IP3
BAI ^D 1	G45IP3
FEZ1	G45IP3
SH3GL3	G45İP3
EF1A	APP1
SETDB1	APP1.
HIP16	APP1
GDF9	APP1
G45IP1	APP1
BAIP1	APP1
HIP5	BAIP3
GIT1	BAIP3
BAIP2	BAIP3

BAIP3

FEZ1	BAIP3
NAG4	BAIP3
SETDB1	BAIP3
HBO1	BAIP3
HIP15	BAIP3
BAIP3	BAIP3
HZFH	BAIP3
PLIP	BAIP3
mHAP1	BAIP3
PiASy	BAIP3
HMP	BAIP3
NAG4	NEFL
HZFH	NEFL
VIM	NEFL
PIASy	NEFL
HMP .	HIP5
PLIP	HIP5
mHAP1	HIP5
HBO1	HIP5
KPNA2	HIP5
VIM .	HIP5
APP1	HIP5
HIP15	HIP5
NAG4	HIP5
GIT1	HIP5
BAIP1 .	. HIP5
FEZ1	HIP5
CGI-74	HIP5
BAIP2	HIP5
ALEX2	ALEX2
PIASy '	MAGEH1
KPNA2	MAGEH1
SETDB1	CA150
LUC7B1	CA150
HZFH ·	CA150
PIASy	CA150
PIASy	hADA3
BAIP1	hADA3
MAGEH1	hADA3
Ku70	hADA3
GIT1	BARD1

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BAIP3 BARD1 SETDB1 BARD1 CA150 BARD1 NAG4 BARD1 HIP15 BARD1 HIP5 BARD1 PTN BARD1 BARD1 FEZ1 **IKAP** BARD1 BAIP1 BARD1 mHAP1 BARD1 HBO1 BARD1 BARD1 BAIP2 PLIP BARD1 **PIASy** BARD1 **HZFH** BARD1 BARD1 ZHX1 SH3GL3 · HDexQ20 HIP13 HDexQ20 CGI-125 HDexQ20 PFN2 HDexQ20 CA150 HDexQ20 HYPA HDexQ20 HP28 HDexQ51 HDexQ51 **HYPA** CA150 HDexQ51 SH3GL3 HDexQ51 HDexQ51 HIP13 HIP15 HDexQ51 PFN2 HDexQ51 CGI-125 HDexQ51 GADD45G LUC7B1 GDF9 GADD45G PTN GADD45G GADD45G BAIP3 G45IP2 GADD45G HIP16 GADD45G GADD45G G45IP3 CGI-125 GADD45G G45IP1 GADD45G HIP5 GADD45G

EF1G	GADD45G
EF1A	GADD45G
PLIP	GADD45G
PIASy .	GADD45G
CGI-74	GADD45G
PTPK	GADD45G
MAP1lc3	PIASy
SUMO-2	PIASy
SUMO-3	PIASy
HYPA	HD1.7
HIP16	HD1.7
DRP-1	HD1.7
HZFH	HD1.7
SH3GL3	HD1.7
HIP13	HD1.7
CGI-125	HD1.7
CA150	HD1.7
HIP11	HD1.7
Ku70	HD1.7
HIP1	HD1.7
IKAP	HD1.7
PFN2	HD1.7
FEZ1	· HD1.7
GIT1	HD1.7
HIP5	HD1.7
PIASy	HD1.7
GIT1	HDd1.0
IKAP	HDd1.0
FEZ1	HDd1.0
PIASy	HDd1.3
IKAP	HDd1.3
HZFH	HDd1.3
Ku70	HDd1.3
PIASy	HIP2
Ku70	CLH-17
HZFH	mp53
ZHX1.	mp53
p53	mp53
PIASy	mp53
PLIP	GAPD

PIASy	IMPD2
EF1G	EF1G
HIP11	EF1G
HZFH	TAL1
ZHX1	TAL1
Ku70	TCPG
PIASy	CLK1
mHAP1	ZNF33B
ZHX1	ZNF33B
HZFH	KPNB1
PIASy	KPNB1
PTN	KPNB1
ALEX2	VIM
SH3GL3	VIM
PIASy	VIM
HIP16	VIM
HBO1	VIM
BAIP1	VIM .
DRP-1	VIM
G45IP1	VIM
MOV34	VIM
VIM	VIM
NEFL	VIM
HSPC232	VIM
SETDB1	VIM
HIP15	HD1.7
HP28	HDexQ20

•
network
interaction
s disease
duntington's
proteins in F
Classification of
Table 7

<u>Q</u>	NAME	FUSION	Tocus ID	ACCESSION	NEO	aa MATCH). FOC
Huntingtin fragments	nents						
HD1.7	huntinatin	OBD	3064	P42858	100	1-506	υ Ż
HDd1.0	huntingtin	OBD	3064	P42858	100	1-320	υ Ž
HDd13	huntinatin	DBD	3064	P42858	100	166-506	υ Ž
HDex020	huntinotin	DBD	3064	P42858	96	1-90	o, Ž
HDexQ51	huntingtin	DBD	3064	P42858	75	1-82	N, C
Transcriptional	Transcriptional control and DNA maintenance						
BARD1	BRCA1 associated ring domain protein 1	080	580	Q99728	66	1-379	Z
CA150	outstive transcription factor CA150	AD, DBD	. 10915	014776	83	299-659	z
GADD45G	growth arrest and DNA damage inducible protein GADD45 gamma	OBO	10912	095257	100	18-159	z
hADA3	ADA3 like protein	OBD	10474	075528	100	235-432	z
HBO1	histone acetyltransferase binding to ORC	AD, DBD ²	11143	095251	100	1-611	z
HYPA	huntingtin interacting protein HYPA/FBP11 (fragment)	AD, DBD	55660	075400	100	8-422	z Ú
HZEH	zinc finder helicase HZFH	AD, DBD	1107	Q9Y410	100	1830-2000	z
IK A D	IKK complex associated protein	AD, DBD ²	8518	095163	9	1207-1332	ς Ż
102 II	ATP denendent DNA helicase II. 70 kDa subunit	AD, DBD1	2547	P12956	100	298-608	z
NAM A	homodomain containing protein NAG4	AD	29117	Q9NPI1	100	94-651	z
PIASY	profein inhibitor of activated STAT protein gamma (PIASv)	AD, DBD	51588	Q8N2W9	100	5-510	υ Ź
153	cellular fumor antiden 553	ΑĐ	7157	P04637	100	1-393	z
p53c	cellular fumor antiden 533 (C-terminus)	ΑĐ	7157	P04637	100	248-393	z
mn53	cellular fumor antiden p53 (mouse)	DBD	7157	P02340	100	73-390	z
20 0	CPI A2 Interacting protein	AD, DBD1	10524	095624	100	5-461	Nd ,
SETUB1	histone-Ivsine N-methyltransferase, H3 lysine-9 specific 4	AD, DBD1	6986	Q15047	100	1023-1291	z
SI 180-2	Inhanition like protein SMT3A (SUMO-2)	. AD	6612	P55854	100	1-103	z Č
2-0M0-3		AD, DBD	6613	P55855	100	1-95	z Ú
ZHX1 ·		AD, DBD	11244	Q9UKY1	100	145-873	z
7815220		DBD	7582	Q8NDW3	100	527-778	z
ZINTOOD	ZIIIC IIIIgal protein coc						

AD, DBD DBD AD						1		
th heavy chain 1 amail dynein light chain (hp28) amail dynein light chain (hp28) agtin associated protein 1 (mouse) aptin interacting protein 1 Ab, DBD ² Ab, DBD APP1	amyloid like protein 1 precursor	AD, DBD	333	P51693	100	243-555	PM, EC	
amal dynein light chain (hp28) and dynein light chain (hp28) agtin associated protein 1 (mouse) agtin interacting protein 1 by DBD² agtin interacting protein 1 by DBD² aptin beta-1 subunit beta-1 subunit beta-1 subunit beta-1 subunit beta-1 subunit anolecular weight neurofilament protein anolecular weight neurofilament protein by DBD² anolecular weight neurofilament protein and lin li by DBD² by DBD² containing GRB2 like protein 3 by DBD² by	CLH-17	clathrin heavy chain 1	DBD	1213	Q00610	10	1-289	PM, V
agin associated protein 1 (mouse) AD, DBD2 AD, DBD3 Illin pherin alpha-2 subunit pherin alpha-2 subunit pherin beta-1 subunit pherin beta-1 subunit pherin beta-1 subunit pherin beta-1 subunit AD, DBD2 AD, DBD3	HP28	axonemal dynein light chain (hp28)	AD	7802	OSBOZe	100	3-258	S
rubule associated protein 1 AD, DBD ² pherin alpha-2 subunit pherin beta-1 subunit pherin precursor (exon 1 included) pherin pherin protein 3 pherin limit (C-terminus) pherin protein 3 pherin pherin pherin pherin 1 (C-terminus) pherin	mHAP1		AD, DBD ²	9001	035668	100	3471	C, EE
hip pherin alpha-2 subunit polerin beta-1 subunit beta-1 subunit beta-1 subunit beta-1 subunit beta-1 subunit beta-1 subunit DBD at DBD and associated proteins 1A/18 light chain 3 AD, DBD and a and a AD,	HP.	huntingtin interacting protein 1	AD, DBD ²	3092	000291	100	245-631	O, GN
Ab, DBD² pherin alpha-2 subunit pherin beta-1 subunit pherin beta-1 subunit pherin beta-1 subunit pherin beta-1 subunit tutbule associated proteins 1A/18 light chain 3 Ab, DBD² Ab,	HMP	mitofilin	· AD, DBD	10989	Q16891	100	212-758	Mit
tubule associated proteins 1A/18 light chain 3 Ab, DBD² Trolecular weight neurofilament protein Trolecular weight neurofilament neurofilamen	KPNA2	karyopherin alpha-2 subunit	AD, DBD ²	3838	P52292	100	141-529	z Ö
tubule associated proteins 1A/1B light chain 3 The cular weight neurofilament protein The cular weight neurofilament protein The part of part of protein 3 The part of p	KPNB1	karyopherin beta-1 subunit	080	3837	014974	100	668-876	z o
Trolecular weight neurofilament protein Trolecular weight neurofilament Trolecular weight neurofilamen	MAPIc3	signaturing accordated anatoine 40/48 light chair 3	AD DRD ²	84557	O9H491	100	58-170	CN. MT
Induction weight reducination process. In II It is to be protein 3 In II In	(כאון האואו)	linch molocular uniaht perindilament profeio	משט מא	4747	081172	100	1-543	N. N.
containing GRB2 like protein 3 build cepeat protein ALEX2 dillo repeat protein ALEX2 build repeat protein ALEX2 dillo repeat protein ALEX2 build repeat protein ALEX2 dulto and elongation protein 2 eta 1 AD, DBD AD	בוצום בוצום	יייייייייייייייייייייייייייייייייייייי	AD, DBD ¹	5217	P35080	9	1-139	Z
containing GRB2 like protein 3 bull of C-terminus) fulful repeat protein ALEX2 full repeat AD, DBD	NTG.	hin precursor (exon	AD, DBD	5764	P21246	8	1-168	PM, EC
ntin (C-terminus) AD AD AD AD AD, DBD AD, DBD In kinase CLK1 AD, DBD AD, DB	SH3G13	SH3 containing GRB2 like protein 3	AD, DBD ²	6457	Q99963	100	3-347	>
dillo repeat protein ALEX2 dillo repeat protein ALEX2 dillo repeat protein ALEX2 dillo repeat protein ALEX2 DBD AD, DBD AD, DBD²			080	7431	P08670	9	1-465	CN, F
dillo repeat protein ALEX2 DBD In kinase CLK1 DBD AD,	VING	vimentin (C-terminus)	AD	7431	P08670	100	189-465	CN, IF
dillo repeat protein ALEX2 In kinase CLK1 In kinase CLK1 In kinase CLK1 In kinase CLK1 In kinase clated protein 1 (C-terminus) In kinase related protein 2cta 1 AD, DBD²	Cell signaling	and fate						
bbD ydropyrimidinase related protein 1 (C-terminus) ydropyrimidinase related protein 1 (C-terminus) bciculation and elongation protein zeta 1 AD, DBD² AD, DBD³	ALEX2	armadillo repeat protein ALEX2	AD, DBD	. 9823	060267	100	127-632	C, PM
ydropyrimidinase related protein 1 (C-terminus) AD, DBD² AD, DBD² AD, DBD³ AD, DBD	C K	protein kinase CLK1	ОВО	1195	P49759	100	209-484	z
sciculation and elongation protein zeta 1 Ab, DBD² Ab, DBD² Ab, DBD² Ab, DBD² 2	DRP-1		AD, DBD¹	1400	Q14194	9	345-572	ပ
Ab, DBD¹ 24 GTPase activating protein GIT1 (9 as insertion included) Ab, DBD² 25 Ab, DBD² 26 Ab, DBD³ 27 Ab, DBD³ 28 Ab, DBD³ 28 Ab, DBD³ 28 Ab, DBD³ 29 Ab, DBD³ 20 Ab, DBD³ 20 Ab, DBD³ 20 Ab, DBD³ 21 Ab, DBD³ 22 Ab, DBD³ 21 Ab, DBD³ 22 Ab, DBD³ 22 Ab, DBD³ 23 Ab, DBD³ 24 Ab, DBD³ 26 Ab, DBD³ 27 Ab, DBD³ 28 Ab, DBD³	FEZ1		AD, DBD ²	9638	Q99689	100	131-392	C, PM
RF GTPase activating protein GIT1 (9 aa insertion included) AD, DBD² Stein-tyrosine phosphatase kappa precursor recraidehyde 3-phosphate dehydrogenase DBD Sisine-5-monophosphate dehydrogenase 2 Insaldolase Internover Insaldolase Instalton factor 1 alpha 1 Insaldon factor 1 gamma AD, DBD Insaldon factor 1 gamma AD, DBD Iquitin conjugating enzyme E2-25 kDa AD, DBD Iquitin conjugating enzyme E2-25 kDa AD, DBD	GDF9	growth/differentiation factor 9	AD, DBD ¹	2661	060383	100	276-454	ပ
oteln-tyrosine phosphatase kappa precursor AD, DBD¹ reraldehyde 3-phosphate dehydrogenase 2 DBD resaldehyde 3-phosphate dehydrogenase 2 DBD resaldelase rad turnover rad turnover ranslation elongation factor 1 alpha 1 AD, DBD ranslation factor 1 gamma (C-terminus) AD republication factor 1 gamma (C-terminus) AD republication factor 1 gamma (C-terminus) AD AD RAD PAD PAD PAD PAD PAD PA	GIT 6	ARF GTPase activating protein GIT1 (9 aa insertion included)	AD, DBD ²	28964	Q9Y2X7	8 6	249-761	PM, V
rceraldehyde 3-phosphate dehydrogenase 2 DBD DBD DBD DBD DBD Insaldolase Interiored a furnover Institution elongation factor 1 alpha 1 AD, DBD Ingation factor 1 gamma (C-terminus) AD Ingation factor 1 gamma	PTPK	protein-tyrosine phosphatase kappa precursor	AD, DBD1	5796	Q15262	100	1227-1439	PM, AJ
rceraldehyde 3-phosphate dehydrogenase 2 DBD Dsine-5'-monophosphate dehydrogenase 2 DBD DBD Insaldolase Interiorer Interi	Cellular metab	olism						
ophosphate dehydrogenase 2 DBD DBD ngation factor 1 alpha 1 AD, DBD AD, DBD tor 1 gamma (C-terminus) gating enzyme E2-25 kDa AD, DBD	GAPD	rceraldehyde 3-phosphate	080	2597	P04406	100	116-334	ပ
ngation factor 1 alpha 1 AD, DBD¹ tor 1 gamma (C-terminus) AD gating enzyme E2-25 kDa AD, DBD	IMPD2	inosine-5'-monophosphate dehydrogenase 2	DBD	3615	P12268	9	34-514	ပ
ngation factor 1 alpha 1 AD, DBD tor 1 gamma (C-terminus) AD gating enzyme E2-25 kDa AD, DBD	TAL1	transaldolase	DBD	6888	P37837	19	3-337	O
translation elongation factor 1 alpha 1 elongation factor 1 gamma c elongation factor 1 gamma (C-terminus) ubiquitin conjugating enzyme E2-25 kDa AD	Protein synthe	sis and turnover						
elongation factor 1 gamma c elongation factor 1 gamma (C-terminus) AD AD ubiquitin conjugating enzyme E2-25 kDa AD AB AD AB	EF1A		AD, DBD1	1915	P04720	100	294-462	C, MT
c elongation factor 1 gamma (C-terminus) AD ubiquitin conjugating enzyme E2-25 kDa DBD MOV34 isolog AD, DBD¹	EF1G	elongation factor 1 gamma	AD, DBD	1937	P26641	19	2-437	C, MT
ubiquitin conjugating enzyme E2-25 kDa DBD AD, DBD1 AO, OV34 isolog	EF1Gc	elongation factor 1 gamma (C-terminus)	AD.	1937	P26641	100	123-437	C, MT
MOV34 isolog AD, DBD1	HIP2	ubiquitin conjugating enzyme E2-25 kDa	OBD	3093	P27924	. 100	1-200	z oʻ
	MOV34	MOV34 isolog	AD, DBD1	10980	015387	92	1-297	z Ú
	TCPG	tein 1. gamma	DBD	7203	P49368	100	252-544	ပ

Uncharacterized proteins	ed proteins						
BAIP1	BARD1 interacting protein 1[similar to RIKEN cDNA 1810018M11]	AD	84289	Q9BS30	100	1-226	S
BAIP2	BARD1 interacting protein 2 [hypothetical protein]	ΑD	84078	Q9H0I6	100	107-684	S
BAID3	RARD1 interacting protein 3 [hypothetical protein]	AD, DBD	55791	Q96HT4	100	152-436	S
C. II. C.	CGL74 moteln	AD	51631	Q9Y383	100	159-270	S
CGI-125	CGI-125 protein	AD	51003	Q9Y3C7	100	1-131	S
G451P1	GADD45G Interacting protein 1[hypothetical protein]	AD, DBD ²	84060	Q9H0V7	100	1-340	S
G45IP2	GADD45G interacting protein 2 [B2 gene partial cDNA, clone B2E]	AD	. 9842	Q9NYA0	100	566-926	S
G451P3	GADD45G Interacting protein 3 [OK/SW-CL.16]	AD, DBD	•	Q8NI70	100	3-134	S
HIPS	huntingtin interacting protein 5 [hypothetical protein KIAA1377]	AD, DBD	57562	Q9P2H0	100	445-988	z, O
HP1	huntinotin interacting protein 11[hypothetical protein]	AD, DBD1	1891	Q96EZ9	100	176-328	S
11013	huntindtin interacting protein 13 [metastasis suppressor protein]	AD, DBD1	9788	Q96RX2	100	512-755	S
HD75	huntinotin interacting protein 15 [similar to KIAA0443 gene product]	AD	114928	Q96D09	9	663-838	S
1012	huntingtin interacting protein 16 Isimilar to KIAA0266 gene product	AD	10813	Q9BVJ6	100	585-771	<u>S</u>
HSPC232	HSPC232	. AD	51535	Q9P0P6	92	1-319	3
11C7B1	putative SR protein LUC7B1 (SR+89)	ΑD	55692	Q9NQ29	66	116-371	ER
MANORH 4	melanoma associated antiden H1	AD, DBD	28986	Q9H213	100	1-219	N
1	modern in the section of the locality of locality of locality of local l	LIACIN CI OI O	Jan Sur	dentity activa	tion domai	n: DBD. D	NA binding

Abbreviations; aa, amino acids; IDEN, identity; LOC, localization; LOCUS ID, NCBI LocusLink Identity, activation domain; DBD, DNA binding domain; DBD, DNA binding domain; DBD, DNA binding horseins; AJ, adherens junctions; C, cytosol; CN, cytoskeleton; EC, extracellular space; EE, early endosomes; ER, endoplasmic reticulum; IF, intermediate filaments; GN, Golgi network; Mit, mitrocholulars, N, nucleus; PM, plasma membrane; pN, perinuclear; UN, unknown; V, vesicles; [], database annotation.

iD	NAME	FUSION /	ACCESSION	IDEN	aa MATCH	LOC
Transcrip	ional control and DNA maintenance	•				
BARD1	BRCA1 associated ring domain protein 1	DBD	Q99728	. 99	1-379	N
CA150	putative transcription factor CA150	. AD	014776	93	299-629	N
Protein sy	nthesis and turnover					
MOV34	MOV34 Isolog	AD, DBD	O15387	95	. 1-297	C, N
Cell signa	ling and fate			•		
GIT1	ARF GTPase activating protein GIT1	AD	Q9Y2X7	98	249-761	PM, V
HSPC232	HSPC232	AD	Q9P0P6	92	1-319	UN
LUC7B1	putative SR protein LUC7B1 (SR+89)	AD	Q9NQ29	99	116-371	ER

Abbreviations: aa, amino acids; IDEN, identity; LOC, localisation; AD, activation domain; DBD, DNA binding domain; AJ, adherens junctions; C, cytosol; CN, cytoskeleton; EC, extracellular space; EE, early endosomes; ER, endoplasmic reticulum; IF, intermediate filaments; GN, Golgi network; Mit, mitochondria; MT, microtubules; N, nucleus; PM, plasma membrane; PN, perinuclear; UN, unknown; V, vesicles; [], database annotation

BAIT

Table 9: New protein-protein interactions found

PREY

SETDB1	SUMO-3
PIASy	SUMO-3
HZFH	SUMO-3
PIASy	· HYPA
HZFH	HYPA
MAP1lc3	HYPA
ZHX1	HYPA
PIASy	HZFH
HZFH	HZFH
GIT1	HZFH
VIM	HZFH
PIASy	ZHX1
HZFH	ZHX1
VIM	ZHX1
FEZ1	HMP
HZFH	HMP
HMP	HMP
PIASy	HMP
HZFH	PTN
HIP15	PTN
PIASy	PTN
PTN	PTN
FEZ1	PTN
KPNA2	G45IP3
GIT1	G45IP3
BAIP1	G45IP3
FEZ1	G45IP3
SH3GL3	G45IP3
EF1A	APP1
SETDB1	APP1
HIP16	APP1
GDF9	APP1
G45IP1	APP1
BAIP1	APP1
HIP5	BAIP3
GIT1	BAIP3
BAIP2	BAIP3
APP1	BAIP3
FEZ1	BAIP3

NAG4	BAIP3
SETDB1	BAIP3
HBO1	BAIP3
HIP15	BAIP3
BAIP3	BAIP3
HZFH	BAIP3
PLIP	BAIP3
mHAP1	BAIP3
PIASy	BAIP3
HMP	BAIP3
NAG4	NEFL
HZFH	NEFL
VIM	NEFL
PIASy	NEFL
HMP	HIP5
PLIP	HIP5
mHAP1 .	HIP5
HBO1	HIP5
KPNA2	HIP5
VIM	HIP5
APP1	HIP5
HIP15	HIP5
NAG4	HIP5
GIT1	HIP5
BAIP1	HIP5
FEZ1	HIP5
CGI-74	HIP5
BAIP2	HIP5
ALEX2	ALEX2
PIASy	MAGEH1
KPNA2	MAGEH1
SETDB1	CA150
LUC7B1	CA150
HZFH	CA150
PIASy	CA150
PIASy	hADA3
BAIP1	hADA3
MAGEH1	hADA3
Ku70	hADA3
GIT1	BARD1
BAIP3	BARD1

SETDB1	BARD1
CA150	BARD1
NAG4	BARD1
HIP15	BARD1
HIP5	BARD1
PTN _.	BARD1
FEZ1	BARD1
IKAP	BARD1
BAIP1	BARD1
mHAP1	BARD1
HBO1	BARD1
BAIP2	BARD1
PLIP	BARD1
PIASy	BARD1
HZFH .	BARD1
ZHX1	BARD1
HIP13	HDexQ20
CGI-125	HDexQ20
PFN2	HDexQ20
HP28	HDexQ51
HIP13	HDexQ51
HIP15	HDexQ51
PFN2	HDexQ51
CGI-125	HDexQ51
LUC7B1	GADD45G
GDF9	GADD45G
PTN	GADD45G
BAIP3	GADD45G
G45IP2	GADD45G
HIP16	GADD45G
G45 P3	GADD45G
CGI-125	GADD45G
G45IP1	GADD45G
HIP5	GADD45G
EF1G	GADD45G
EF1A	GADD45G
PLIP .	GADD45G
PIASy	GADD45G
CGI-74	GADD45G
PTPK .	GADD45G
MAP1Ic3	PIASy

SUMO-2	PIASy
SUMO-3	PIASy
HIP16	HD1.7
DRP-1	HD1.7
HZFH	HD1.7
HIP13	HD1.7
CGI-125	HD1.7
HIP11	HD1.7
Ku70	HD1.7
IKAP	HD1.7
PFN2	HD1.7
FEZ1	HD1.7
GIT1	HD1.7
HIP5	, HD1.7
PIASy	HD1.7
GIT1	HDd1.0
IKAP	HDd1.0
FEZ1	HDd1.0
PIASy	HDd1.3
IKAP	HDd1.3
HZFH	HDd1.3
Ku70	HDd1.3
PIASy	HIP2
Ku70	CLH-17
HZFH	mp53
ZHX1	mp53
p53	mp53
PIASy	mp53
PLIP	GAPD
PIASy	IMPD2
EF1G	EF1G
HIP11	EF1G
HZFH	TAL1
ZHX1	、TAL1
Ku70	TCPG
PIASy	CLK1
mHAP1	ZNF33B
ZHX1	ZNF33B
HZFH	KPNB1
PIASy	KPNB1
PTN	KPNB1

ALEX2 VIM SH3GL3 VIM PIASy VIM HIP16 VIM HZFH VIM HBO1 VIM BAIP1 VIM DRP-1 VIM G45IP1 VIM MOV34 VIM VIM VIM NEFL VIM HSPC232 VIM SETDB1 VIM HIP15 HD1.7 HP28 HDexQ20

	entary Table 1: List of DBD proteins for			
ID	NAME	ACCESSION	aa MATCH	PPIs
BARD1	BRCA1 associated ring domain protein 1	Q99728	1-379	3
CLH-17	clathrin heavy chain 1	Q00610	1-289	1
CLK1	protein kinase CLK1	P49759	209-484	1
GADD45G	growth arrest and DNA-damage-Inducible protein GADD45 gamma	O95257	18-159	6
hADA3 ·	ADA3 like protein	075528	235-432	1.
HD1.7	huntingtin	P42858	1-506	5
HDd1.0	huntingtin	P42858	1-320	1
HDd1.3	huntingtin	P42858	166-506	2
HDexQ20	huntingtin	P42858	1-90	3
HDexQ51	huntingtin	P42858	1-82	4
HIP2	ubiquitin conjugating enzyme E2-25 kDa	P27924	1-200	1
MPD2	inosine-5'-monophosphate dehydrogenase 2	P12268	34-514	1
KPNB1	karyopherin beta-1 subunit	Q14974	668-876	1
mp53	cellular tumor antigen p53 (mouse)	P02340	73-390	2
TAL1	transaldolase	P37837	3-337	1
TCPG	T-complex protein 1, gamma subunit	P49368	252-544	1
VIM	vimentin	P08670	1-465	6
ZNF33B	zinc finger protein 33b	Q8NDW3	527-778	1
14-3-3	14-3-3 protein epsilon	P42655	93-255	AA
DNAJ	DnaJ homolog subfamily A member 1	P31689	113-379	AA
HD513Q68	huntingtin	P42858	1-513	AA
HIP1	huntingtin interacting protein 1	O00291	245-631	AA
mAP2A1	œadaptin A (mouse)	P17426	697-971	AA
mAP2A2	œadaptin C (mouse)	P17427	697-938	AA
mHAP	huntingtin associated protein 1 (mouse)	O35668	3-471	AA
RFA	replication protein A 70 kDa DNA-binding subunit	P27694	262-616	AA
SH3GL3	SH3 containing GRB2 like protein 3	Q99963	3-347	AA
ZFR	ZNF259	075312	29-460	AA
ACTG1	gammaactin	P02571	182-375	• ′
ALBU	serum albumin precursor	P02768	1-249	-
ALDA	fructose-bisphosphate aldolase A	P04075	1-363	-
AMPL	cytosol aminopeptidase	· P28838	46-487	-
ARF4L	ADP-ribosylation factor-like protein 4L	P49703	33-201	-
ASNS	glutamine-dependent asparagine synthetase	P08243	318-560	-
BCK	creatine kinase, B chain	P12277	92-381	-
CLH-17	clathrin heavy chain 1	Q00610	1165-1671	-
GAPDH	glyceraldehyde 3-phosphate dehydrogenase	P04406	1-334	-
HD-CT	huntingtin	P42858	2721-3144	-
LDHB .	L-lactate dehydrogenase b chain	P07195	96-333	-
MDHM	malate dehydrogenase, mitochondrial precursor	P40926	1-338	-
MOV34	MOV34 isolog	015387	76-297	-
NSFL1C	p97 cofactor p47	Q9UNZ2	201-370	-
PEBP	phosphatidylethanolamine-binding protein	P30086	1-186	-
PHGDH	D-3-phosphoglycerate dehydrogenase	043175	1-553	-
PLD2	phospholipase D2	O14939	168-336	-
TIP49	49 kDa TBP-interacting protein	Q9Y265	1-456	_
TRFE	serotransferrin precursor	P02787	213-698	-
TUBA1	alpha-tubulin 1	P05209	1-451	-
TUBB4	tubulin beta-4 chain	Q13509	113-450	-
UBC1	polyubiquitin C	Q9UEF2	1-685	

Abbreviations: aa, amino acids; DBD, DNA binding domain; PPIs, protein-protein interactions; AA, autoactivation of reporter gene.

Suppler	mentary Table 2: Subcloned DBD proteins for 2 ⁿ	Tourid of library screens
Prey	Reason for selection	PPIs
HIP5	huntingtin interacting protein verified by in vitro binding assay	8
PIASy	huntingtin interacting protein verified by <i>in vitro</i> binding assay	3 ·
CA150	huntingtin interacting protein, literature verified interaction [Holbert S. et al. Proc. Natl Acad. Sci. USA 98, 1811-1816 (2001)]	1
EF1G	part of ternary complex with EF1A, which is found in htt aggregates [Vanwetswinkel S. et al. J Biol. Chem. 278, 43443-51 (2003)]	1 .
HYPA	huntingtin interacting protein, literature verified interaction [Faber, P.W. et al. Hum. Mol. Genet.9, 1463-1474 (1998)]	1
FEZ1	huntingtin interacting protein verified by in vitro binding assay	AA
GIT1	huntingtin interacting protein verified by in vitro binding assay	AA
EF1A	htt aggregate-interacting protein [Mitsui K. et al. J. Neurosci.22,9267-9277 (2002)]	-
HIP11	huntingtin interacting protein verified by in vitro binding assay	÷
NEFL.	vimentin interacting protein, literature verified interaction [Carpenter, D.A. & lp; W. <i>J. Cell. Sci.</i> 10, 2493-2498 (1996)]	-
p53	huntingtin interacting protein, literature verified interaction [Steffan, J.S. <i>et al. Proc. Natl. Acad. Sci. USA</i> 97 , 6763-8 (2000)]	
PLIP	BARD1 interacting protein, literature verified interaction [Dechend, R. et al. Oncogene 18, 3316-3323 (1999)]	•

Abbreviations: DBD, DNA binding domain; PPIs, protein-protein interactions; AA, autoactivation of reporter gene.

Supplementary Table 3: Reported interactions in Huntington's disease network

Denorted	interactions.	found
Renorted	interactions.	KHIHKI

Protein A	Protein B	Literature
CA150	HD1.7 HDexQ20 HDexQ51	Holbert S. et al. Proc. Natl Acad. Sci. USA 98, 1811-1816 (2001). The Gin-Ala repeat transcriptional activator CA150 interacts with huntingtin: neuropathologic and genetic evidence for a role in Huntington's disease pathogenesis.
HYPA	HD1.7 HDexQ20 HDexQ51	Faber, P.W. et al. Hum. Mol. Genet.9, 1463-1474 (1998). Huntingtin Interacts with a family of WW domain proteins.
HIP1	HD1.7	Wanker, E.E. et al. Hum. Mol. Genet.3, 487-495 (1997). HIP-I: a huntingtin interacting protein isolated by the yeast two-hybrid system.
SH3GL3	HD1.7 HDexQ20 HDexQ51	Sittler, A. et al. Mol. Cell ⁴ , 427-436 (1998). SH3GL3 associates with the Huntingtin exon 1 protein and promotes the formation of polygln-containing protein aggregates.
PIASy	mp53	Nelson, V., Davis, G.E. & Maxwell, S.A. <i>Apoptosis</i> 3, 221-234 (2001). A putative protein inhibitor of activated STAT (PIASy) interacts with p53 and inhibits p53-mediated transactivation but not apoptosis.
p53	mp53	Chene, P. Oncogene 20, 2611-2617 (2001). The role of tetramerization in p53 function. Leblanc V. et al. Anal Biochem 308, 247-54 (2002). Homogeneous timeresolved fluorescence assay for identifying p53 interactions with its protein partners, directly in a cellular extract.
PLIP	BARD1	Dechend, R. et al. Oncogene 18, 3316-3323 (1999). The Bcl-3 oncoprotein acts as a bridging factor between NF-kappaB/Rel and nuclear co-regulators.
SUMO-2	PIASy	Sachdev, S. et al. Genes Dev.15, 3088- 3103 (2001). PIASy, a nuclear matrix-associated SUMO E3 ligase, represses LEF1 activity by sequestration into nuclear bodies.
SUMO-3	PIASy	Sachdev, S. et al. Genes Dev.15, 3088- 3103 (2001). PIASy, a nuclear matrix-associated SUMO E3 ligase, represses LEF1 activity by sequestration into nuclear bodies.
EF1G	EF1G	Mansilla, F. et al. Biochem. J.365, 669-676 (2002). Mapping the human translation elongation factor eEF1H complex using the yeast two-hybrid system.
NEFL VIMc	VIM	Carpenter, D.A. & Ip, W. J. Cell. Sci.10, 2493-2498 (1996). Neurofilament triplet protein interactions: evidence for the preferred formation of NFL containing dimers and a putative function for the end domains.

Reported interactions, not found

Protein A	Protein B	Literature
HAP1	HDexQ20 HDexQ51	Li, S.H. et al. J. Biol. Chem. 273, 19220-19227 (1998) A human HAP1 homologue. Cloning, expression, and interaction with huntinglin.
	HDexCol	Li, S.H. et al. <i>J. Neurosci.</i> 18 , 1261-1269. (1998) Interaction of huntingtin-associated protein with dynactin P150Glued.
HIP1	CLH-17	Henry, K.R. et al. Mol. Bio.I Celis, 2607-2625 (2002). Scd5p and clathrin function are important for cortical actin organization, endocytosis, and localization of sla2p in yeast. [interlogs paper]
•		Metzler, M. et al. J. Biol. Chem. 276, 39271-39276 (2001). HIP1 functions in clathrin-mediated endocytosis through binding to clathrin and adaptor protein 2.
		Waeiter, S. et al. Hum. Mol. Genet.10, 1807-1817 (2001). The huntingtin interacting protein HIP1 is a clathrin and alpha-adaptin-binding protein involved in receptor-mediated endocytosis.
p53	HDexQ20 HDexQ51	Steffan, J.S. et al. Proc. Natl. Acad. Sci. USA 97, 6763-6768 (2000). The Huntington's disease protein interacts with p53 and CREB-binding protein and represses transcription.
p53	hADA3	Wang, T. et al. EMBO J.20, 6404-6413 (2001). hADA3 is required for p53 activity.
p53	BARD1	Irminger-Finger, I. et al. Mol. Cell6, 1255-1266 (2001). Identification of BARD1 as mediator between proapoptotic stress and p53-dependent apoptosis.
KPNA2	KPNB1	Chook, Y.M. & Blobel, G. Curr. Opin. Struct. Biol.6, 703-715 (2001). Karyopherins and nuclear import.

Supple	mentary Table 4: Reported huntingtin inter	doung protont	
D .	NAME	LOCUS ID	PubMed ID
ranscript	ional control and DNA maintenance		
CA150	transcription elongation regulator 1 (TCERG1)	10915	11172033
CREB1	cAMP responsive element binding protein 1	1385	8643525
CREBBP	CREB binding protein (Rubinstein-Taybl syndrome)	1387	10823891
CTBP1	C-terminal binding protein 1	1487	11739372
-IYPA	formin binding protein 3 (FNBP3)	55660	9700202 .
-IYPB	huntingtin interacting protein B	29072	9700202
HYPC	huntingtin interacting protein C	25766	9700202
ICOR1	nuclear receptor co-repressor 1	9611	10441327
	nuclear factor of kappa light polypeptide gene enhancer in		40070454
VFKB1	B-cells 1 (p105)	4790	12379151
PQBP1	polyglutamine binding protein 1	10084	10332029
REST	RE1-silencing transcription factor	5978	1288172
SAP30	sin3-associated polypeptide, 30kDa	8819	10823891;10441327
SP1	Sp1 transcription factor	6667	11988536
TAF4	TAF4 RNA polymerase II	6874	11988536
ГВР	TATA box binding protein	6908	10410676
TP53	tumor protein p53 (Li-Fraumeni syndrome)	7157	10823891
Celiular o	rganization and protein transport		
AP2A2	adaptor-related protein complex 2, alpha 2 subunit	_. 161	9700202
DLG4	discs, large homolog 4 (Drosophila) (PSD95)	1742	11319238
HAP1	huntingtin-associated protein 1 (neuroan 1)	9001	9668110;9454836
HIP1	huntingtin interacting protein 1	3092	9147654
HIP14	huntingtin interacting protein 14	23390	9700202;12393793
OPTN '	optineurin (FIP2)	10133	9700202;11137014
PACSIN1	protein kinase C and casein kinase substrate in neurons 1	29993	12354780
SH3GL3	SH3-domain GRB2-like 3	6457	9809064
SYMPK	symplekin `	8189	9700202
TUBG1	tubulin, gamma 1	7283	11870213
Cell signa	ling and fate		
GRAP	GRB2-related adaptor protein	10750	8612237
GRB2	growth factor receptor-bound protein 2	2885	9079622
ITPR1	inositol 1,4,5-triphosphate receptor, type 1	3708	12873381
MAP3K10	mitogen-activated protein kinase kinase kinase 10	4294	10801775
PDE1A	phosphodiesterase 1A, calmodulin -dependent	5136	8643525
RASA1	RAS p21 protein activator (GTPase activating protein) 1	5921	8612237; 9079622
TGM2	transglutaminase 2	7052	11442349
TRIP10	thyroid hormone receptor interactor 10	9322	12604778
	netabolism		
CBS	cystathionine-beta-synthase	875	9466992; 10434301;10823891
GAPD	glyceraldehyde-3-phosphate dehydrogenase	2597	8612237
TPH1	tryptophan hydroxylase 1	7166	12354289
	ynthesis and tumover		
HIP2	huntingtin interacting protein 2	3093	8702625; 9700202
	terized proteins		
Oncharac HYPE	huntingtin interacting protein E	11153	9700202
HYPK	huntingtin interacting protein HYPK	25764	9700202
HYPM	huntingtin interacting protein HYPM	25763	9700202
MAGEA3		4102	9700202

Abbreviations: ID, interacting protein gene symbol; LOCUS ID, NCBI LocusLink Identity; Pubmed ID, NCBI PubMed publication index; Reported htt interactors are presented according to databases: MINT, HPRD, BIND; Li & Li, *Trends Genet.* (2004), **20** 146-152 and Harjes & Wanker, *Trends. Biochem. Sci.* (2003), **28**, 425-433.

Supplementary Table & Protein-protein interactions of the extended HD network

Number	ID 1	LOCUSID 1	ID 2	LOCUSID 2	Reference	Number	ID 1	LOCUSID 1	ID 2	LOCUSID 2	Reference
1	ABL1	25	CBL	867	literature	63	BRCA 1	672	RBBP4	5928	literature
ż	ABL1	25	PXN	5829	literature	64	BRCA1	672	RELA	5970	literature
3	ALEX2	9823	ALEX2	9823	this study	65	CA150	10915	LUC7B1	55692	this study
4	ALK	238	SHC1	6464	literature	66	CA150	10915	PIASy	51588	this study
5	AP2A2	161	SHC1	6464	literature	67	CBL	867	SRC	6714	literature
. 6	APP1	333	EF1A	1915	this study ·	68	CBL.	867	VAV1	7409	literature
7	APP1	333	BAIP1	84289	this study	69	CBL	867	SH3KBP1	30011	literature
8	APP1	333	GDF9	2661	this study	70	CBL	867	LAT	27040	literature
9	APP1	333	SETBD1	9869	this study	71	CBL.	867	SHC1	6464	literature
10	APP1	333	HIP16	10813	this study	72	CBL	867	PIK3R1	5295	literature
11	APP1	333	BAIP3	55791	this study	73	CBL	867	PLCG1	5335	literature
12	APP1	333	HIP5	57562	this study	74	CBL	867	FYN	2534	literature
13	APP1	333	G45IP1	84060	this study	75	CBL	867	PTK2B	2185	literature
14	AR	367	EP300	2033	literature	76	CBL	867	EGFR	1956	literature
15	AR	367	ESR1	2099	literature	77	CDC2	983	PCNA	5111	literature
16	AR	367	RELA	5970	literature	78	CDC2	983	FYN	2534	literature
17	AR	367	BRCA1	672	literature	79	CGI-74	5163 1	HIP5	57562	this study
18	AR	367	HDAC1	3065	literature	80	CHUK	1147	IKBKB	3551	literature
19	AR	367	NCOA1	8648	literature	· 81	CLH-17	1213	HGS	9146	literature
20	AR	367	JUN	3725	literature	82	CLH-17	1213	Ku70	2547	this study
21	AR	367	NCOA3	8202	literature	83	CLK1	1195	PIASy	51588	this study
22	AR	367	STAT3	6774	literature	84	CREB1	1385	BRCA1	672	literature
23	AR	367	NR3C1	2908	literature	85	CREB1	1385	NR3C1	2908	literature
24	BAIP1	84289	G45IP3		this study	86	CREBBP	1387	MSX1	4487 -	literature
25	BAIP3	55791	BAIP2	84078	this study	87	CREBBP	· 1387	RELA	5970	literature
26	BAIP3	55791	HIP15	114928	this study	88	CREBBP	1387	RBBP4	5928	literature
27	BAIP3	55791	BAIP3	55791	this study	89	CREBBF	1387	PTMA	5757	literature
28	BAIP3	55791	HIP5	57562	this study.	90	CREBBF	1387	PPARG	5468	literature
29	BARD1	580	PLIP	10524	this study	91	CREBBF	1387	PML	537,1	literature
30	BARD1	580	ZHX1	11244	this study	92	CREBBF	1387	MYOD1	4654	literature
31	BARD1	580	POU2F1	5451 `	literature	93	CREBBF	1387	JUN	3725	literature
32	BARD1	580	BRCA1	672	literature	94	CREBBF	1387	HNF4A	3172	literature
33	BARD1	580	CA150	10915	this study	95	CREBBF	1387	NR3C1	2908	 literature
34	BARD1	580	GIT1	28964	this study	. 96	CREBBE		EVI1	2122	literature
35	BARD1	580	IKAP	8518	this study	97	CREBBF	1387	KLF5	688	literature
36	BARD1	580	HBO1	11143	this study	98	CREBBE		SRC	6714	literature
37	BARD1	580	CDC2	983	literature	99	CREBBF		BCL3	602	literature
38	BARD1	580	NAG4	29117	this study	100	CREBBF	1387	TP53	7157	literature
39	BARD1	580	BAIP2	84078	this study	101	CREBBF	1387	BRCA1	· 672	literature
40	BARD1	580	PIASy	51588	this study	102	CREBBF	1387	WT1	7490	literature
41	BARD1	580	BAIP3	55791	this study	103	CREBBE	2 1387	NCOA3	8202	literature
42	BARD1	580	HIP5	57562	this study	104	CREBBF	1387	NCOA1	8648	literature
43	BARD1	580	SETBD1	9869	this study	105	CREBBE	1387	KHDRBS ¹		literature
44	BARD1	. 580	BCL3	602	literature	106	CREBBE		HIPK2	28996	literature
45 .	BARD1	580	HAP1	9001	this study	107	CREBBE	P 1387	SREBF2		literature
46	BARD1	580	PTN	5764	this study	108	CREBBE		AR	367	literature
47	BARD1	580	HZFH	1107	this study	109	CTBP1	1487	HDAC2	3066	literature
48	BARD1	580	HIP15	114928	this study	110 .	CTBP1	1487	ZNFN1A		literature
49	BARD1	580	BAIP1	84289	this study	111	. CTBP1	1487	HDAC1	3065	literature
50	BARD1	580	FEZ1	9638	this study	112	CTBP1	1487	EVI1	2122	literature
51	BCL3	602	FYN	2534	literature	113	CTBP1	1487	BRCA1	672	literature
52	BCL3	602	RXRA	6256	literature	114	DLG4	1742	HGS	9146	literature
. 53	BCL3	602	JUN	3725	literature	115	DLG4	1742	ÉΥΝ	2534	literature
54	BCL3	602	SHC1	6464	literature	116	DLG4	1742	PRKCA	5578.	literature
55	BRCA1		HDAC2		literature	117	DLG4	1742	DNCL1	8655	literature
56	BRCA1		EP300	2033	literature	118	DLG4	1742	ERBB2	2064	literature
57	BRCA1		ESR1	2099	literature	119	DRP-1	1400	Huntingti		this study
58	BRCA1		CDC2	983	literature	120	DRP-1	1400	VIM	7431	this study
59	BRCA1		HDAC1		literature	121	EF1A	1915	GADD450		this study
60	BRCA1		STAT3		literature	122	EF1A	1915	PLCG1	5335	literature
61	BRCA1		JUN	3725	literature	123	EF1G	1937	EF1G	1937	this study
62	BRCA1	672	MYC	4609	literature	124	EF1G	1937	GADD45	G 10912	this study

Number	ID 1	LOCUSID 1	ID 2	OCUSID 2	Reference	Number	ID 1	LOCUSID 1	ID 2	LOCUSID 2	Reference
125	EGFR	· 1956	SRC	6714	literature	190	GIT1	28964	BAIP3	55791	this study
126	EGFR	1956	PTK2	5747	literature	191	GIT1	28964	G451P3		this study
127	EGFR	1956	PLCG1	5335	literature	192	GIT1	28964	HIP5	57562	this study
128	EGFR	1956	PIK3R1	5295	literature	193	GIT1	28964	PXN	5829	literature
129	EGFR	1956	ERBB2	2064	literature	194	GIT1	28964	PTK2	5747	literature
130	EGFR	1956	PDGFRB	5159	literature	195	GRAP	10750	EPOR	2057	literature
131	EGFR	1956	PTK2B	2185	literature	196	GRAP	10750	TNFSF6	356	literature
132	EGFR	1956	ESR1	2099	literature	197	GRAP	10750	KIT	3815	literature
133	EGFR	1956	SHC1	6464	literature	198	GRAP	10750	SOS1	6654	literature
134	EGFR	1956	SOS1	6654	literature	199	GRAP	10750	LAT	27040	literature
135	EP300	2033	ING1	3621	literature	200	GRB2	2885	TP73L PLCG1	8626 5335	literature literature
136	EP300	2033	NCOA1	8648	literature	201	GRB2	2885			
137	EP300	2033	HNF4A	3172	literature	202	GRB2	· 2885 2885	PTK2 SHC1	5747 6464	literature literature
138	EP300	2033	MDM2	4193	literature	203 204	GRB2 GRB2	2885	SOS1	6654	literature
139	EP300	2033	PCNA PTMA	5111 5757	literature	204	GRB2	2885	LAT	27040	literature
140	EP300	2033 2033	RELA	5970	literature literature	206	GRB2	2885	SRC	6714	literature
141 142	EP300 EP300	2033	STAT3	6774	literature	207	GRB2	2885	WAS	7454	literature
143	EP300	2033	ESR1	2099	literature	208	GRB2	2885	WASL	8976	literature
144	EPOR -	2057	KIT	3815	literature	209	GRB2	2885	KHDRBS1	10657	literature
145	EPOR	2057	SHC1	6464	literature	210	GRB2	2885	SH3KBP1	30011	literature
146	EPOR	2057	VAV1	7409	literature	211	GRB2	2885	PIK3R1	5295	literature
147	EPOR	2057	PIK3R1	5295	literature	212	GRB2	2885	RASA1	5921	literature
148	ERBB2	2064	PTK2	5747	literature	213	GRB2	2885	VAV1	7409	literature
149	ERBB2	2064	SHC1	6464	literature	214	GRB2	2885	EGFR	. 1956	literature
150	ERBB2	2064	PTK2B	2185	literature	215	GRB2	2885	ABL1	25	literature
151	ERBB2	2064	." SOS1	6654	literature	216	GRB2	2885	TNFSF6	356	literature
152	ESR1	2099	JUN	3725	literature	217	GRB2	2885	PDGFRB	5159	literature
153	ESR1	2099	MDM2	4193	titerature	218	GRB2	2885	DNM1	1759	literature
154	ESR1	2099	PIK3R1	5295	literature	219	GRB2	2885	EPOR	2057	literature
155	ESR1	2099	SHC1	6464	literature	220	GRB2	2885	ERBB2	2064	literature
156	ESR1	2099	NCOA3	8202	literature	221	GRB2	2885	PTK2B		literature
157	ESR1	2099	NCOA1	8648	literature	222	GRB2	2885	HRAS	3265 3815	literature literature
158	EVI1	2122	HDAC1	3065	literature	223	GRB2 GRB2	2885 2885	KIT CBL	867	literature
159	FEZ1	9638	HMP	10989	this study	224			FGFR1	2260	literature
160	FEZ1	9638	BAIP3	55791	this study	225 · 226	GRB2 hADA3	2885 10474	EP300	2033	literature
161	FEZ1	9638	HIP5	57562 	this study this study	227	hADA3	10474	TP53	7157	literature
162	FEZ1	9638	G45IP3	6464	literature	228	hADA3	10474	BAIP1	84289	this study
163 164	FGFR1 FYN	2260 2534	SHC1 VAV1	7409	literature	000	hADA3	10474	PIASy	51588	this study
165	FYN	2534	SHC1	6464	literature	230	hADA3	10474	MAGEH1	28986	this study
166	FYN	2534	KHDRBS1	10657	literature	231	hADA3	10474	ESR1	2099	literature
167	FYN	2534	WAS	7454	literature	232	HAP1	9001	BAIP3	55791	this study
168	FYN	2534	PDGFRB	5159	literature	233	HAP1	9001	HGS	9146	literature
169	FYN	2534	PIK3R1	5295	literature	234	HAP1	9001	HIP5	57562	this study
170	FYN	2534	PLCG1	5335	literature	235	HBO1	11143	MCM2	4171	literature
171	FYN	2534	PXN	5829	literature	236	HBO1	11143	HIP5	57562	this study
172	FYN	2534	PTK2	5747	literature	237	HBO1	11143	BAIP3	55791	this study
173	G45IP2	9842	GADD45G	10912	this study	238	HBO1	11143	AR	367	literature
174	GADD450	3 10912	G45IP1	84060	this study	239	HDAC1	3065	PML	5371	literature
175	GADD450		HIP5	57562	this study	240	HDAC1	3065	RELA	5970	literature
176	GADD450	3 10912	LUC7B1	55692	this study	241	HDAC1	3065	PTMA	5757	literature
177	GADD450		RXRA	6256	literature	242	HDAC1	3065	PHB	5245	literature
178	GADD450		BAIP3	55791	this study	243	HDAC1	3065	MYOD1	4654 5111	literature literature
179	GADD450		PIASy	51588	this study	244	HDAC1	3065	PCNA		
180	GADD450		G45IP3	 E468	this study	245	HDAC1	3065 3065	RBBP4 ING1	5928 3621	literature literature
181	GADD450		PPARG	5468	literature	246	HDAC1 HDAC1	3065	HDAC2	3066	literature
182	GADD450		PCNA ESP1	5111 2099	literature literature	247 248	HDAC1		PTMA	5757	literature
183	GADD450		ESR1 CDC2	983	literature	249	HDAC2		RBBP4	5928	literature
184 185	GADD450		CGF125	51003	this study	250	HIP11	1891	. EF1G	1937	this study
186	GADD45		CGF123	51631	this study	251	HIP11	1891	Huntingtir		this study
. 187	GAPD	2597	DNCL1	8655	literature	252	HIP16	10813	GADD450		this study
188		2597	PLIP	10524	this study		HIP2	3093	PIASy	51588	this study
189	GDF9	2661	GADD45G		this study	254	HIP2	3093	TP53	7157	literature
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Number	ID 1 HIP5	LOCUSID 1	ID 2 BAIP2	LOCUSID 2 84078	Reference this study	Number 320	· ID 1 Huntingtin	LOCUSID 3064	I ID 2 OPTN	10133	Reference literature
255 256	HIP5	57562 57562	BAIP1	84289	this study	321.	HYPA	55660	MAP1Ic3	84557	this study
257	HIP5	57562	HIP15	114928	this study	322	HZFH	1107	SUMO-3	6613	this study
258	HMP	10989	PIASy	51588	this study	323	HZFH	1107	MIV	7431	this study
259	HMP	10989	HIP5	57562	this study	324	HZFH	1107	HZFH	1107	this study
260	HMP	10989	HMP	10989	this study	325	HZFH	1107	, Huntingtin	3064	this study
261	HMP	10989	BAIP3	55791	this study	326	HZFH	1107	BAIP3	55791	this study
262	HNF4A	3172	NCOA3	8202	literature	327	HZFH	1107	HYPA PIASy	55660 51588	this study this study
263	HNF4A	3172	SRC	6714 6721	literature . literature	328 329	HZFH HZFH	1107 1107	GIT1	28964	this study
264 265	HNF4A HRAS	3172 . 3265	SREBF2 SOS1	6654	literature	330	HZFH	1107	ZHX1	11244	this study
266	HRAS	3265	VAV1	7409	literature	331	HZFH	1107	NEFL.	4747	this study
267	HRAS	3265	PIK3R1	5295	literature	332	HZFH	1107	CA150	10915	this study
268	HRAS	3265	MAPK8	5599	literature	333	HZFH	1107	TP53	7157	this study
269	Huntingtir		TUBG1	7283	literature	334	HZFH	1107	PTN	5764	this study
270	Huntingtin		RASA1	5921	literature	335	HZFH	1107	KPNB1	3837	this study
271	Huntingtir		HYPA	55660	this study	336	HZFH	1107	TAL1 HMP	6888 10989	this study this study
272	Huntingtin		GRB2 HIP1	2885 3092	literature this study	337 338	HZFH IKAP	1107 8518	CHUK	1147	literature
273 274	Huntingtin		· HIP2	3092	literature	339	IKAP	8518	IKBKB	3551	literature
275	Huntingtir Huntingtir		MPR1	3708	literature	340	IKAP	8518	MAPK8	5599	literature
276	Huntingtir		REST	5978	literature	341	IMPD2	3615	PIASy	51588	this study
277	Huntingtin		MAGEA3		literature	342	ING1	3621	PCNA	5111	literature
278	Huntingtir	3064	SH3GL3	6457	this study	343	ING1	3621	RBBP4	5928	literature
279	Huntingtir		HAP1	9001	literature	344	JUN	3725	STAT3	6774	literature
280	Huntingtir		SYMPK	8189	literature	345	JUN	3725	RELA MYOD1	5970 4654	literature literature
281	Huntingtir		TBP	6908	literature	346 347	JUN	3725 3725	NCOA1	8648	literature
282	Huntingtir		SP1 NFKB1	6667 47 9 0	literature literature	348	JUN	3725	MAPK8	5599	literature
283 284	Huntingtir Huntingtir		PDE1A	5136	· literature	349	KIT	. 3815	PIK3R1	5295	literature
285	Huntingtir		TAF4	6874	literature	350	KIT	3815	PLCG1	5335	literature
286	Huntingtin		GAPD	2597	literature	351	KPNA2	3838	G45IP3		. this study
287	Huntingtin		TPH1	7166	literature	352	KPNA2	3838	MAGEH1	28986	this study
288	Huntingtir		TP53	7157	literature	353	KPNA2	3838	DD5	51366	literature
289	Huntingtin		TGM2	7052	literature	354	KPNA2	3838	RELA	5970 5757	literature
290	Huntingtin		MAP3K10		literature	355	KPNA2 KPNA2	3838 3838	PTMA TP53	5757 7157	literature literature
291	Huntingtin		SAP30 CREB1	8819 1385	literature literature	356 357	KPNA2	3838	HIP5	57562	this study
292 293	Huntingtii Huntingtii		HIP15	114928	this study	358	KPNB1	3837	TP53	7157	literature
293 294	Huntingti		PIASy	51588	this study	359	KPNB1	3837	PIASy	51588	this study
295	Huntingti		CGF125	51003	this study	360	KPNB1	3837	PTN	5764	this study
296	Huntingti		GIT1	28964	this study	361	KPNB1	3837	DD5	51366	literature
297	Huntingti	n 3064	HIP16	10813	this study	362	KPNB1	3837	PTMA	5757	literature
298	Huntingti		HIP13	9788	this study	363	KPNB1	3837	FGFR1	2260	literature
299	Huntingti		FEZ1	9638	this study	364 365	Ku70 Ku70	2547 2547	hada 3 TCPG	10474 7203	this study this study
300	Huntingti		ikap HP28	8518 7802	this study this study	366	Ku70	2547	Huntingtin		this study
301 302	Hunfingtii Hunfingtii		PFN2	5217	this study	367	Ku70	2547	EGFR	1956	literature
303	Huntingti		HYPK	25764	literature	368	Ku70	2547	PCNA	5111	literature
304	Huntingti		DLG4	1742	literature	369	Ku70	2547	MAPK8	5599	literature
305	Huntingti		HYPE	11153	literature	370	Ku70	2547	VAV1	7409	literature
306	Huntingti	n 3064	CREBBF		literature	371	Ku70	2547	PTTG1	9232	literature
307	Huntingti		CA150	10915	this study	372	Ku70	2547	WRN	7486	literature
308	Huntingti		NCOR1	9611	literature	373	Ku70	2547	ABL1	25 51588	literature this study
309	Huntingti		PACSIN1		literature	374 375	MAGEH1 MAP3K10	28986 4294	PIASy PHB	5245	literature
310 311	Huntingti Huntingti		HYPB PQBP1	29072 10084	literature literature	376	MAP3K10		RACGAP1		literature
311	Huntingti		CTBP1	1487	· literature	377	MDM2	4193	PML	5371	literature
313	Huntingti		GRAP	10750	literature	378	MEN1	4221	RELA	5970	literature
314	Huntingti		TRIP10	9322	literature	379	MYC	.4609	MAPK8	5599	literature
315	Huntingti	n 3064	HYPC	25766	literature	380	MYC	4609	RELA	5970	literature
. 316	Huntingti		HIP14	23390	literature	381	MYOD1	4654	RXRA	6256	literature
317	Huntingti		HYPM	25763	literature		MYOD1	4654	STAT3	6774 57562	literature this study
318	Huntingti		AP2A2	161 975	literature	383 384	NAG4 NAG4	29117 29117	HIP5 BAIP3	55791	this study
319	Huntingti	n 3064	CBS	875	literature	304	IVAG4	28111	UMI-3	00131	uno saudy

Number	ID 1	LOCUSID 1	ID 2 L	OCUSID 2	Reference	Number	ID1	LOCUSID 1	ID 2	LOCUSID 2	Reference
Number 385	NCOR1	9611	PML	5371	literature	450	РТК2В	2185	PIK3R1	5295	literature
386	NCOR1	9611	ESR1	2099	literature	451	PTK2B	. 2185	PXN	5829	literature
387	NCOR1	9611	PHB	5245	literature	452	PTK2B	2185	FYN	. 2534	literature
388	NCOR1	· 9611	PTMA	5757	literature	453	PTK2B	2185	SRC	6714	literature
389	NCOR1	9611	NCOA3	8202	literature	454	PTK2B	2185	VAV1	7409	literature
390	NCOR1	9611	AR	367	literature	455	PTN	5764	GADD45G		this study
391	NCOR1	9611	NR3C1	2908	literature	456	PTN	5764	FEZ1	9638	this study
392	NEFL	.4747	TSC1	7248	literature	457	PTN	5764	PTN	5764	this study
393	NEFL	4747	PRKCL1	5585	literature	458	PTN	5764	ALK	238	literature
394	NEFL	4747	PIASy .	51588	this study	459	PTN	5764	PIASy	51588	this study
395	NEFL	4747	VIM	7431	this study	460	PTN	5764 5706	HIP15 GADD45G	114928 10912	this study this study
396	NEFL	4747	NAG4	29117	this study .	461	PTPK	5796	SRC	6714	literature
397	NFKB1	4790	CHUK	1147	literature	462 463	PXN RASA1	5829 5921	PTK2B	2185	literature
398	NFKB1	4790 4790	AR KLF5	367 688	literature literature	464	RASA1	5921	PIK3R1	5295	literature
399 400	NFKB1 NFKB1	4790 4790	NR3C1	2908	literature	465	RASA1	5921	PDGFRB	5159	literature
401	NFKB1	4790	MEN1	4221	literature	466	RASA1	5921	HRAS	3265	literature
402	NFKB1	4790	IKBKB	3551	literature	467	RASA1	5921	FYN	2534	literature
403	NFKB1	4790	BRCA1	672	literature	468	RASA1	5921	PXN	5829	literature
404	NFKB1	4790	STAT3	6774	literature	469	RASA1	5921	ALK	238	literature
405	NR3C1	2908	NCOA1	8648	literature	470	RASA1	5921	SRC	6714	literature
406	NR3C1	2908	RELA	5970	literature	471	RASA1	5921	KHDRBS1		literature
407	NR3C1	2908	MDM2	4193	literature	472	RELA	5970	STAT3	6774	literature
408	NR3C1	2908	STAT3	6774	literature	473	RXRA	6256	NCOA3	8202	literature
409	NR3C1	2908	JUN	3725	literature	474	SAP30	8819	ING1	3621	literature
410	PACSIN1		WASL	8976	literature	475	SAP30	8819	HCFC1	3054	literature
411	PACSIN1		DNM1	1759	literature	476	SAP30	8819	HDAC1	3065	literature
412	PCNA	5111	PTMA	5757	literature	477	SAP30	8819	HDAC2	3066	literature
413	PCNA	5111	WRN	7486	literature	478	SAP30	. 8819	RBBP4	5928	literature literature
414	PDGFRB		PLCG1	5335	literature	479	SAP30	8819	NCOR1 CA150	9611 _. 10915	this study
415	PDGFRB		SHC1	6464	literature	480	SETBD1	9869 9869	BAIP3	55791	this study
416	PDGFRB		PIK3R1	5295	literature	481 482	SETBD1 SH3GL3	6457	VIM	7431.	this study
417	PDGFRB		PTK2	5747 84557	literature this study	483	SH3GL3	6457	G45IP3		this study
418	PIASy PIASy	51588 51588	MAP1lc3 BAIP3	55791	this study	484	SH3GL3	6457	CBL	867	literature
419 420	PIASy	. 51588	HYPA	55660	this study	485	SH3GL3	6457	SH3KBP1		literature
421	PIK3R1	5295	SHC1	6464	literature	486	SOS1	6654	LAT	27040	literature
422	PIK3R1	5295	SRC	6714	literature	487	SOS1	6654	SH3KBP1	30011	literature
423	PIK3R1	5295	VAV1	7409	literature	488	SP1	- 6667	HNF4A	3172	literature
424	PIK3R1	5295	WAS	7454	literature	489	SP1	6667	HCFC1	3054	literature
425	PIK3R1	5295	HGS	. 9146	literature	490	SP1	6667	BRCA1	672	literature
426	PIK3R1	5295	KHDRBS1	10657	literature	491	SP1	6667	HDAC1	3065	literature
427	PIK3R1	5295	LAT	27040	literature	492	SP1	6667	HDAC2	3066	literature
428	PIK3R1	5295	PTK2	5747	literature	493	SP1	6667	JUN	3725	literature
429	PLCG1	5335	LAT	27040	literature	494	SP1	6667	MSX1	4487	literature
430	PLCG1	5335	WAS	7454	literature	495	SP1	6667	MYC	4609	literature
431	PLCG1	5335	SOS1	6654	literature	496	SP1	6667	MYOD1	4654	literature
432	PLCG1	5335	SRC	6714	literature	497	SP1	6667	PML POU2F1	5371 5451	literature literature
433	PLCG1	5335	VAV1	7409	literature	498 499	SP1 SP1	6667 6667	RBBP4	5928	literature
434	PLCG1	5335	KHDRBS1	10657	literature				RXRA	6256	literature
435	PLIP	10524	BCL3	602	literature	500 501	SP1 SP1	6667 6667	SHC1	6464	· literature
436	PLIP PLIP	10524 10524	AR STAT3	367 6774	literature literature	502	SP1	6667	SREBF2		literature
437 438	PLIP	10524	GADD45G	10912		503	SP1	6667	KLF4	9314	literature
439	PLIP	10524	BAIP3	55791	this study	504	SP1	6667	TP53	7157	literature
440	PLIP	10524	HIP5	57562	this study	505	SRC	6714	KHDRBS		literature
441	PML	5371	RELA	5970	literature	506	SRC	6714	WAS	7454	literature
442	PPARG		RXRA	6256	literature	507	SRC	6714	STAT3	6774	literature
443	PPARG		NCOA1	8648	literature	508	STAT3	6774	NCOA1	8648	literature
444	PQBP1		AF	367	literature	509	STAT3	6774	KHDRBS		literature
445	PRKCA		YWHAZ	7534	literature	510	SUMO-2	6612	PIASy	51588	this study
446	PTK2	5747	PXN	5829	literature	511	SUMO-3		PIASy	51588	this study
447	PTK2	5747	SHC1	6464	literature	512	SUMO-3		PML	5371	literature
448	PTK2	5747	SRC	6714	literature	513	SUMO-3		SETBD1		this study
449	PTK2B	2185	SHC1	6464	literature-	514	TAF1B	9014	TAF1A	9015	literature

Number	ID 1	LOCUSID 1	ID 2	LOCUSID 2	Reference
515	TAF1C	9013	TAF1B	9014	literature ·
516	TAF1C	9013	TAF1A	9015	literature
517	TAL1	6888	ZHX1	11244	this study
518	TBP	6908	TAF1B	9014	literature
519	TBP	6908	MSX1	4487	literature
520	TBP	6908 6908	HMGB1 NR3C1	3146 2908	literature literature
521 522	TBP TBP	6908	MCM2	4171	literature
522 523	TBP	6908	MDM2	4193	literature
523 524	TBP	6908	MYC	4609	literature
525	TBP	6908	RXRA	6256	literature
526	TBP	6908	NCOA3	8202	literature
527	TBP	6908	BCL3	602	literature
528	TBP	6908	TAF1C	9013	literature
529	TBP	6908	TP53	7157	literature
530	TBP	6908	TAF1A	9015	literature
531	TBP	6908	ZNFN1A1		literature
532	TBP	6908	JUN NCOA1	3725 8648	literature literature
533 534	TBP TNFSF6	6908 356	FYN	2534	literature
534 535	TNFSF6	356 356	SRC	6714	literature
536	TP53	7157	HMGB1	3146	literature
537	TP53	7157	YWHAZ	7534	literature
538	TP53	7157	NR3C1	2908	literature
. 539	TP53	7157	HNF4A	3172	literature
540	TP53	7157	ING1	3621	literature
541	TP53	7157	PIASy	51588	this study
542	TP53	7157	PML	5371	literature
543	TP53	7157	EP300	2033	literature
544	TP53	7157	MAPK8 CHUK	5599 1147	literature literature
545 546	TP53 TP53	7157 7157	WT1	7490 ·	literature
546 547	TP53	. 7157	MDM2	4193	literature
548	TP53	7157	TP73L	8626	literature
549	TP53	7157	TAF1C	9013	literature
550	TP53	7157	TAF1B	9014	literature
551	TP53	7157 .	TAF1A	9015	literature
552	TP53	7157	PTTG1	9232	literature
553	TP53	7157	KLF4	9314	literature
554	TP53	7157	HIPK2	28996	literature
555	TP53	7157 7457	WRN BRCA1	7486 672	literature literature
556 557	TP53 TP53	7157 7157	ABL1	25	literature
558	TP53	7157	TP53	7157	this study
559	TP53	7157	ZHX1	11244	this study
560	TP53	7157	PRKCA	5578	literature
561	TP53	7157	CDC2	983	literature
562	TP73L	8626	HIPK2	28996	literature
563	TRIP10	.9322	RXRA	6256	literature
564	TRIP10	9322	WAS	7454	literature
565	TSC1	7248	YWHAZ	7534	literature
566 567	TUBG1 TUBG1	7283 - 7283	PIK3R1 BRCA1	5295 672	literature literature
568	TUBG1	7283	PXN	5829	literature
569	TUBG1	7283	RACGAP'		literature
570	VAV1	7409	LAT	27040	literature
571	VIM	7431	MEN1	4221	literature
572	VIM	7431	PRKCL1	5585	literature
573	VIM .	7431	TSC1	7248	literature
574	VIM	7431	DNCL1	8655	literature
575	VIM	7431	HIP16	10813	this study
576	VIM	7431	YWHAZ		literature
577 579	VIM	7431 7431	VIM SETBD1	7431 9869	this study this study
578 579	VIM VIM	7431 7431	MOV34	10980	this study
318	AllAf	1401	1110 104	,0300	a ao olaay

Number	ID 1	LOCUSID 1	ID 2	LOCUSID 2	Reference
580	VIM	7431	HBO1	11143	this study
-581	VIM	7431	ZHX1	11244	this study
582	VIM	7431	HSPC232	51535	this study
583	VIM	7431	PIASy	51588	this study
584	VIM	7431	HIP5	57562	this study
585	VIM	7431	G45IP1	84060	this study
586	VIM	7431	BAIP1	84289	this study
587	VIM.	7431	ALEX2	9823	this study
588	ZHX1	11244	HYPA	55660	this study
589	ZHX1	11244	PIASy	51588	this study
590	ZNF33B	7558	HAP1	9001	this study
591	ZNF33B	7558	ZHX1	11244	this study
					

Abbreviations: ID, interacting protein gene symbol; LOCUS ID, NCBI LocusLink Identity. The presented list of protein-protein interactions is computed from databases: MINT, HPRD, BIND; Li & Li, Trends Genet. (2004), 20, 146-152 and Harjes & Wanker, Trends. Biochem. Sci. (2003), 28, 425-433.

The figures show:

Figure 1 Identification of two-hybrid interactions connected to HD. a, Schematic representation of the screening strategy. b, Identification of interactions by systematic interaction mating. Upper panel: Selection of diploid yeast clones by transfer on minimal medium lacking leucine and tryptophan (SDII). Lower panel: Two-hybrid selection of interactions on minimal medium lacking leucine, tryptophan, histidine and uracil (SDIV) after 5 days of growth at 30°C. The prey proteins HP28 (A5), SH3GL3 (A7), CA150 (B9), HIP15 (B10), PFN2 (B11), HIP13 (C1), CGI125 (C12) and HYPA (D1) were identified as HDexQ51 interactors.

Figure 2 Protein interaction network for Huntington's disease. a, Matrix of 117 two-hybrid interactions between 21 bait and 49 prey proteins. b, Yeast two-hybrid interactions depicted as network using the software Pivot 1.0. In total, 96 interactions and 61 distinct proteins are depicted. In addition, dimers of EF1G, VIM and p53 are shown.

Figure 3 Systematic validation of two-hybrid interactions by *in vitro* binding experiments. GST-fusion proteins (baits) immobilised on glutathione agarose beads were incubated with COS1 cell extracts containing HA-tagged prey proteins. After extensive washing of the beads, bound proteins were eluted and analysed by SDS-PAGE and immunoblotting using anti-HA antibody.

Figure 4 Identification of network proteins stimulating htt aggregation. a, Filter retardation assay. Protein extracts were prepared from HEK293 cells coexpressing HD169Q68 and network proteins as indicated. The aggregated proteins retained on the filter were detected with anti-htt antibody (CAG53b) and anti-GIT1 antibody. b, Coimmunoprecipitation of HD510Q68 and GIT1 from COS1 cell extracts. Extracts were incubated with anti-GIT1 or preimmune serum. Immunoprecipitated material was analysed by immunoblotting using htt- antibody 4C8 and anti-HA antibody. c, Coimmunoprecipitation of htt and GIT1 from human brain extracts. Protein complexes containing GIT1 were pulled-down with increasing amounts of anti-htt antibodies, but not with corresponding preimmune sera. d, Analysis of subcellular localisation of HD510Q68 and GIT1 by immunofluorescence microscopy. COS1 cells were transfected with the indicated constructs and immunolabled with 4C8 anti-htt

antibody coupled to Cy3-conjugated antibody (red) and with anti-HA antibody coupled to FITC-conjugated antibody (green). Nuclei were counterstained with Hoechst (blue). Colocalisation of HD510Q68 and GIT1 is illustrated by yellow colour of the insoluble aggregates. Scale bars, $10~\mu m$.

Figure 5 Detection of GIT1 in brains of R6/1 transgenic mice and HD patients. a, Sections of striatum and cortex of R6/1 mice brains labelled with anti-GIT1 and anti-htt (EM48) antisera. Arrows point to nuclear inclusions. b, Inclusions in cortex of HD patients are labelled with anti-htt (2B4) and anti-GIT1 antibodies. Arrows indicate neuronal inclusions, recognized by anti-htt (2B4) and anti-GIT1 antibodies. Scale bars, $20~\mu m$. c, Colocalisation of GIT1 and htt in the cortex of HD patients detected by immunofluorescence microscopy.

Figure 6 Amino acid sequence of the interacting proteins of the PPI of huntingtin.

Figure 7 Identification of Y2H interactions connected to HD. A, The screening strategy. B, Identification of interactions by systematic interaction mating. Upper panel: Selection of diploid yeast clones on SDII minimal medium. Lower panel: Two-hybrid selection of interactions on SDIV minimal medium. The prey proteins HP28 (A5), SH3GL3 (A7), CA150 (B9), HIP15 (B10), PFN2 (B11), HIP13 (C1), CGI125 (C12), and HYPA (D1) were identified as HDexQ51 interactors.

Figure 8 A protein interaction network for Huntington's disease. A, Matrix of 186 Y2H interactions between 35 bait and 51 prey proteins. Interactions reported previously (30), or verified in pull down assays (35) are indicated. B, A comprehensive PPI network for htt. Y2H interactors identified in this study (red diamonds), previously published interactors (blue squares), interactors identified from databases HRPD, MINT and BIND, bridging any two proteins in the extended network (green triangles, Suppl. Table 5). Htt interactors previously reported and found in our screens (CA150, HYPA, HIP1, and SH3GL3), depicted as red squares.

Figure 9 Validation of Y2H interactions by *in vitro* binding experiments. GST-fusion proteins immobilized on glutathione agarose beads were incubated with COS-1 cell extracts containing HA-tagged proteins. After extensive washing, pulled proteins were eluted and analyzed by SDS-PAGE and immunoblotting using anti-htt 4C8 or anti-HA antibodies.

Figure 10 GIT1 enhances and is critical for htt aggregation. A, Filter retardation assay for the identification of GIT1 as a promoter of htt aggregation. 48 h post transfection, protein extracts were prepared from HEK293 cells coexpressing HD169Q68 and GIT1-CT (aa 249-770). Aggregated proteins retained on the filter were detected with anti-htt (CAG53b) or anti-C-GIT1 antibody. B, Effect of full-length GIT1 on HD169Q68 aggregation analyzed by the filter retardation assay. C, Analysis of HD169Q68 aggregation in cells overexpressing GIT1-CT by indirect immunofluorescence microscopy. a, HD169Q68 (red). b, GIT1-CT (green). c, Colocalization of GIT1 with the endosomal marker EEA1 is indicated in yellow. d-f, Colocalization of HD169Q68 (red) and GIT1-CT (green) in COS-1 cells. D, Silencing of endogenous GIT1 expression. HEK293 cells transfected with the siRNA-GIT1 were analyzed after 48 h by immunoblotting using anti-C-GIT1 and anti-GAPDH antibodies. E, Silencing of endogenous GIT1 prevents the accumulation of insoluble htt aggregates. siRNA-GIT1 treated and untreated cells expressing HD169Q68 were analyzed 72 h post transfection by filtration.

Figure 11 Verification of the htt–GIT1 interaction. A, Coimmunoprecipitation of HD510Q68 and HA-GIT1-CT from COS-1 cell extracts using anti-C-GIT1 antibody. Immunoprecipitated material was analyzed by immunoblotting, using the anti-HA 12CA5 antibody detecting recombinant GIT1 (upper blot) and the htt-4C8 antibody (lower blot). B, Coimmunoprecipitation of htt and GIT1 from human brain extracts. C, Subcellular localization of GIT1 and htt in differentiated PC12 cells (a-c) and SH-SY5Y cells (d-f) by confocal immunofluorescence microscopy. Colocalization of htt and GIT1 shown in yellow (panel c and f). Arrow points to cytoplasmic structures recognized by both antibodies. In addition, specific GIT1 labeling was detected at the tip of neurite-like extensions in adhesion foci (arrowheads). Scale bars, 10 μm.

Figure 12 Detection of GIT1 in brains of transgenic mice and HD patients. A, Sections of striatum and cortex of R6/1 mice brain labeled with anti-C-GIT1 and anti-htt EM48 antibodies. Arrows point to nuclear inclusions. B, Neuronal inclusions (arrows) in cortex of HD patients recognized by anti-htt 2B4 and anti-C-GIT1 antibodies. Scale bars, 20 μm. C, Colocalization of GIT1 and htt in the cortex of HD

patients, detected by immunofluorescence microscopy. D, Detection of N-terminally truncated GIT1 degradation products in HD patient brain cortex.

Figure 13 Specificity of GIT1 antibodies. A, Scheme indicating the regions of GIT1, which were used for the production of antibodies. NT-GIT1 antibody recognizes the N-terminal part (aa 1-100), C-GIT1 the central part (aa 368-587) and CT-GIT1 the C-terminal part (aa 664-770) of GIT1. B, Analysis of the specificity of the GIT1 antibodies. All three antibodies specifically recognize GIT1, but not the homologous protein GIT2 (Premont et al., 2000). After expression of full length HAGIT1 and HAGIT2 15 μg of total COS-1 cell extract was subjected to SDS-PAGE. Immunoblotting was performed with anti-NT-GIT1 (1:500), anti-C-GIT1 (1:500) and anti-CT-GIT1 (1:500) antibodies. Expression of HA-GIT1 and HA-GIT2 was detected with an anti-HA antibody (1:1000).

The examples illustrate the invention:

PART I: ESTABLISHING THE PROTEIN-INTERACTION NETWORK OF HUNTINGTIN

Examples 1: Particular methods and material used in the Examples

• Antibodies, strains and plasmids

A polyclonal antibody (pAb) against GIT1 was generated by injection of affinity purified His₆-tagged GIT1 (residues 368-587) into a rabbit. The htt-specific pAb CAG53b and HD1 were described ^{13,14}. Commercially available antibodies were anti-GST pAb (Amersham Pharmacia), anti-GIT1 pAb (Santa Cruz Biotechnology), anti-HA monoclonal antibody 12CA5 (mAb) (Roche Diagnostics), anti-htt pAb EM48 ⁴⁷, anti-htt mAb 2B4 ⁴⁸ and anti-htt mAb 4C8 (Chemicon). As secondary antibodies for immunofluorescence microscopy Cy3- and FITC-conjugated IgGs (Jackson ImmunoResearch) were used. The yeast strains used as two-hybrid reporters were L40ccua [MATa his3Δ200 trp1-901 leu2-3,112 LYS2::(lexAop)₄-HIS3 ura3::(lexAop)₈-lacZ ADE2::(lexAop)₈- URA3 GAL4 gal80 can1 cyh2] and L40ccα [MATα his3Δ200 trp1-910 leu2-3,112 ade2 LYS2::(lexAop)₄-HIS3 URA3::(lexAop)₈-lacZ GAL4 gal80 can1 cyh2]. Both strains are derivatives of L40c ¹⁷. Plasmids pHD510Q17 and pHD510Q68 were generated by insertion of fragments coding for HD510Q17 and HD510Q68 into pcDNA-I (Invitrogen). pHD169Q68 was derived from pHD510Q68 by deletion of the Xhol- Xhol fragment encoding aa 170-510 of human htt.

Library screening

Plasmids encoding bait proteins were transformed into the strain L40ccua, tested for the absence of reporter gene activity and cotransformed with a human fetal brain cDNA library (Clontech). For each transformation 1 x 10⁶ independent transformants were plated onto minimal medium lacking tryptophan, leucine, histidine and uracil (SDIV medium) and incubated at 30°C for 5 to 10 days. Clones were picked into microtitre plates using a picking robot and grown over night in liquid minimal medium lacking tryptophan and leucine (SDII medium). Then, they were spotted onto nylon or nitrocellulose membranes placed on SDIV medium plates. After incubation for 4 days

membranes were subjected to a β -galactosidase (β -GAL) assay. Plasmids were prepared from positive clones and characterised by restriction analyses and sequencing. For retransformation assays plasmids encoding bait and prey proteins were cotransformed in the yeast strain L40ccua and plated onto SDIV medium.

Array mating screen

Plasmids encoding bait and prey proteins were transformed into strains L40ccua and L40ccα, respectively. L40ccα clones were arrayed in 96-well microtitre plates and mixed with a single L40ccua clone for interaction mating. Diploid cells were transferred by a robot (Beckman, Biomek® 2000) onto YPD medium plates and, after incubation for 24 h at 30°C, onto SDII medium plates for additional 72 h at 30°C. For two-hybrid selection diploid cells were transferred onto SDIV medium plates with and without nylon or nitrocellulose membranes and incubated for 5 days at 30°C. The nylon or nitrocellulose membranes were subjected to the β-GAL assay. Positive clones were verified by cotransformation assays using plasmids encoding respective bait and prey proteins.

Protein expression and verification assays

For verification experiments cDNA fragments encoding baits and preys were subcloned into pGEX derivatives (Stratagene) or pTL-HA ¹⁸. GST fusion proteins were expressed in *E. coli* BL21-codon PlusTM RP (Stratagene) and affinity purified on glutathione agarose beads (Sigma) using standard protocols ¹⁷. COS1 cells were transfected with mammalian expression plasmids and lysed as described ¹⁸. For *in vitro* binding assays, 30 µg of GST or GST fusion protein were immobilized on glutathione agarose beads and incubated with 500 µg protein extract prepared from COS1 cells expressing a HA-tagged fusion protein for 2 h at 4°C in binding buffer [50 mM HEPES pH 7.4, 150 mM NaCl, 10% glycerol, 1 % NP-40, 1 mM EDTA, 20 mM NaF, 1 mM DTT, 0.1 % Triton X-100, protease inhibitors (Roche Diagnostics)]. After centrifugation and extensive washing of the beads bound proteins were eluted and analysed by SDS-PAGE and Western blotting. Coimmunoprecipitation experiments were performed as described by Sittler *et al.* ¹⁸. For immunofluorescence microscopy COS1 cells were grown on cover slips and cotransfected with pcDNA-HD510Q68 and pTL-HA-GIT1. 40 h post transfection cells were fixed with 2% paraformaldehyde.

Standard protocols for staining with appropriate primary and secondary antibodies were used ¹⁸.

• Filter Retardation Assay

HEK293 cells coexpressing HD169Q68 and GIT1, PIASy, HIP5, HP28, PFN2, FEZ1 or BARD1 were harvested 48 h post transfection. Cells were lysed as described ¹⁸ and boiled in 2% SDS, 100 mM DTT for 5 min. Aliquots containing 50, 25 and 12.5 µg of total protein were used for filtration on a cellulose acetate membrane ¹⁴. SDS-resistant aggregates were detected using anti-CAG53b or anti-GIT1 antibodies.

Immunocytochemistry

Mice were deeply anaesthetised and perfused through the left cardiac ventricle with 4% paraformaldehyde in 0.1 M phosphate buffer. Brains were removed and postfixed overnight in 4% paraformaldehyde. Sections were processed for immunocytochemistry as described ⁴⁷. pAb EM48 (1:1000) and affinity purified anti-GIT1 pAb (1:100) were used as primary antibodies.

Six human HD and 5 control brains were used in this study. Two HD cases were classified as grade 3 and four cases as grade 4 of neuropathological severity. For immunolabelling standard protocols were used ⁴⁸. 2B4 mAb (1:200) and affinity purified GIT1 pAb (1:50) were used as primary antibodies.

Example 2: Two-hybrid screens and data management

To generate a PPI network for HD we used a combination of library and matrix yeast two-hybrid screens (Fig. 1a). First, 50 selected cDNAs encoding proteins potentially involved in HD including 10 different htt fragments were cloned into a DNA binding domain vector for expression of LexA fusion proteins (baits). The resulting plasmids were introduced into yeast strain L40ccua, which carries three reporter genes, *HIS3*, *URA3* and *lacZ*, for two-hybrid interaction analyses. Forty baits did not activate the reporters by themselves and were used individually for cotransformation screening of a human fetal brain cDNA library expressing GAL4 activation domain hybrids (preys). In each screen, 1 x 10⁶ auxotrophic transformants were tested on selective plates, and 1-50 positive colonies were typically obtained. Restriction analyses and

sequencing identified preys that together with their respective baits repeatedly activated the reporter genes. Starting with 40 baits in the first round of cotransformation screens we identified 34 PPIs for 10 baits (Table 1).

In the second round of screens, 12 cDNA fragments encoding preys identified in the first screen were subcloned into a DNA binding domain vector. The resulting baits were tested for autoactivation and 10 were screened against a human fetal brain cDNA library. Four of the 10 proteins revealed additional 13 PPIs.

Finally, an array mating screen was performed to connect all baits and preys identified in the transformation screens. For this assay, MATα yeast cultures were transformed with plasmids encoding prey proteins and arrayed in 96-well microtitre plates for interaction mating with individual MATa strains expressing bait proteins. Using this strategy each bait was individually tested for interaction with every prey in the array. Diploid yeast clones, formed by mating on YPD plates, were selected on agar SDII plates, and further transferred by a spotting robot on SDIV plates to select for Y2H interactions (Fig. 1b). We examined 3500 pairwise combinations of baits and preys in the mating assay and identified additional 70 PPIs. These interactions could be confirmed in cotransformation assays (Table 5).

Table 5:

Summary of two-hybrid screens

	baits	preys	baits yielding	interactions	
Screen	screened	screened	interactions	identified	
1st transfr mation screen	40	4x10 ⁷	10	34	
2nd transformation screen	10	1x10 ⁷	4	13	
Array mating screen	50	70	. 21	70	

Thus, the combination of cDNA library and array mating screens proved powerful in establishing a highly connected PPI network linked to htt.

Sequence analysis of the cDNAs encoding bait and prey proteins revealed ORFs ranging from 82 to 728 amino acids in size (Table 2). In a systematic Blast search 60 out of the 67 proteins identified were identical to a SwissProt or TrEMBL protein entry (http://us.expasy.org/sprot/). The remaining 7 proteins showed 75-99 % identity to its best fit and either contained single amino acid substitutions, variable polyQ lengths or small regions of sequence variation. Uncharacterised proteins were named according to their interaction partners. Each ORF was further examined for consensus protein domains using the FprintScan, HMMPfam, HMMSmart, ProfileScan, and BlastProDom programs providing useful hints to protein function. For example, the protein BAIP1 (BARD1 interacting protein 1) possesses a Zn-finger-like PHD finger that is believed to be important for chromatin-mediated transcriptional regulation. Similarly, domain searches for BAIP2 (BARD1 interacting protein 2) revealed a BTB/POZ domain, a motif found in developmentally regulated zinc finger proteins of the Kelch family of actin-associated proteins. Thus, BAIP2 could potentially mediate the association of BARD1 with the actin cytoskeleton.

Example 3: Analysis and functional assignment of the two-hybrid data

Our two-hybrid screens identified a total of 117 PPIs between 70 protein fragments. As a result of the iterative two-hybrid strategy all interactions could be depicted in a single large network. The number of interactions identified for each bait varied from 1 to 18, with each protein having 1.6 interaction partners on average. In order to display the PPI data, both matrix and network representations were used (Fig. 2). The matrix shows, in addition to the two-hybrid interactions, previously reported interactions and interactions verified by independent methods (Fig. 2a). In comparison, the network view allows to immediately recognize local PPI patterns and paths connecting two proteins in the network (Fig. 2b). Interestingly, proteins such as htt, BARD1, GADD45G, HIP5, PIASy or VIM interact with more than 11 other proteins forming nodes within the HD network, while 30 proteins have only one interaction partner and thus are located at the periphery of the network (Fig. 2b). Indeed, all other proteins a e embedded in many bi-fan motifs and multiple circular interaction clusters that have been interpreted to be an indication for biological relevance ^{11,19}. Schwikowski et al. ²⁰ defined network proteins, which are separated

by no more than two other proteins, as being part of a functional cluster. In this respect all proteins in our network form a functional cluster with htt.

We assigned a subcellular localisation to each protein by examining various sources of literature and based on available experimental data we grouped the proteins into six broad functional categories (Fig. 2a, Table 2).

Eighteen proteins in the HD network are involved in transcriptional regulation or DNA maintenance (Fig. 2a). The second largest group, 14 proteins, includes mainly cytoskeletal and transport proteins. We assigned 5 proteins to cellular signalling and fate, another 4 proteins to protein synthesis and turnover, and 3 proteins to cellular metabolism. Being part of 41 interactions, 16 proteins of unknown function were identified.

For the analysis of htt PPIs, as much as 40 out of 117 interactions (34,2%) included a htt fragment (Fig. 2a). In total, 19 different htt interacting partners from various functional groups were detected, 4 proteins had been previously described and 6 involved proteins of unknown function. Surprisingly, most htt partners (6) are involved in transcriptional regulation and DNA maintenance, but others function in cell organization and transport (4), cellular signalling (2), or cellular metabolism (1), suggesting that htt functions in different subcellular processes.

The current hypothesis that htt has a function in transcriptional regulation is inferred from its interactions with transcriptional activators, coactivators or repressors ²¹. In agreement with previous reports, binding of htt to CA150 ²² and HYPA ²³ has been detected in our screens. In addition, new connections to nuclear proteins such as SETBD1, PLIP and HBO1 were found. These multidomain proteins act on histones and are known modulators of chromatin structure and gene expression. Similarly, the zinc finger bromo domain containing proteins BARD1, NAG4, HZFH, ZHX1, ZNF33B play a role in transcriptional control. The protein IKAP directly interacts with htt and was recently shown to be part of a complex regulating RNA polymerase II activity ²⁴. Htt also interacts with PIASy, which inhibits transcription factor STAT-mediated gene activation ²⁵. PIASy functions as SUMO E3 ligase for the Wnt-responsive transcription factor LEF1, inhibiting its activity via sumoylation ²⁶. This suggests that PIASy catalysed sumoylation of transcription factors could represent a general

mechanism in repression of gene expression. The binding of PIASy to htt indicates that htt may itself be a substrate for sumoylation. Alternatively, it could influence the sumovlation of other transcription factors. Thus, our data extend the nuclear role of htt and provide additional leads for its involvement in transcriptional regulation.

Another large group of htt interactors identified here are proteins that function in cellular organization and vesicle transport. We report a new interaction between htt and dynein light chain (HP28), a component of the dynein/dynactin motor protein complex. Interestingly, the p150 Glued subunit of dynactin is linked to the htt-associated protein HAP1. 16,27. Our observation that htt directly binds to HP28 underscores the potential scaffolding role of htt/HAP1 in dynein/dynactin driven retrograde vesicle transport along microtubules in axons.

The htt interacting protein HIP1 anchors clathrin-coated vesicles to the cytoskeleton via its actin-binding domain, a link crucial for synaptic vesicle endocytosis 28. Here, a new PPI between htt and profilin II (PFN2) 29 was detected. PFN2, a protein enriched in neurons, modulates actin polymerization in vitro and is involved in endocytosis via association with scaffolding proteins 29. The htt-PFN2 connection adds support to a potential role of htt in modulation of both actin polymerization and vesicle transport processes.

Currently, for the function of 6 htt interactors, including HIP5, no genetic or biochemical evidence is available (Table 2). We found that HIP5 binds to htt as well as to karyopherin α (KPNA2). KPNA2 serves as an adapter for karyopherin β (KPNB1), which transports NLS-tagged proteins into the nucleus 30. Thus, HIP5 might take this route to the nucleus. Interestingly, HEAT or armadillo (ARM) repeats, forming α -helical structures in KPNA2 and KPNB1 are also present in htt 31 . Therefore, the complexes between KPNA2 and HIP5 as well as between htt and HIP5 could be similar in terms of protein structure. It is tempting to further speculate that htt participates in nucleocytoplasmic transport.

Example 3: Verification of PPIs

Comparison with literature-cited interactions revealed that more than 80% of the twohybrid interactions identified here are novel. For all network bait and prey proteins only 24 PPIs have been reported previously using two-hybrid methods,

81

coimmunoprecipitations or affinity chromatography-based techniques; 18 of these were confirmed in our Y2H assays (Fig 2a, Table 2). Failure to detect interactions may result from the high stringency of our particular two-hybrid system. However, in most cases the occurrence of false negatives can be explained by the lack of essential domains in one of the protein fragments used. For example, an interaction between p53 and hADA3 has been described ³², with the first 214 amino acids of hADA3 being essential for this interaction. It escaped our two-hybrid analysis, because a C-terminal hADA3 fragment (amino acids 235-432) was used. For the same reason, an interaction between p53 and BARD1 or between KPNA2 and KPNB1 was not observed.

Beside false negatives, the two-hybrid assay is also prone to create false positive results ⁹. Addressing this issue we performed a series of pull-down and overlay assays and thereby confirmed several of the two-hybrid PPIs independently. Proteins were expressed as GST-fusions in *E. coli* and as HA-fusions in COS1 cells. After immobilization of the GST-fusion protein to beads or nitrocellulose membranes the respective partner was affinity-purified from a COS1 cell extract and binding was detected by immunoblotting. Using these assays, 22 physical interactions, central to the HD network, were verified (Fig. 2a). The results of some *in vitro* GST pull-down assays are shown in Fig. 3. For example HD510Q17 interacts with HIP1, GIT1, PIASy, FEZ1 and HIP11, and HIP5 binds to HD510Q68, GIT1, HBO1, PLIP and FEZ1 (Fig. 3). In total, 35 two-hybrid interactions were verified independently either in previous studies or by our *in vitro* binding assays (Fig. 2a).

Example 4: GIT1 promotes htt aggregation in vivo

The formation of insoluble polyQ-containing protein aggregates is a pathological hallmark of HD. Several lines of evidence link htt aggregation to disease progression and the development of motor symptoms. We screened network proteins for their potential to enhance htt aggregation in a cell-based aggregation assay ¹⁴. In this assay, formation of SDS-insoluble htt aggregates in mammalian cells, that have been cotransfected with constructs encoding an N-terminal htt fragment with 68 glutamines (HD169Q68) and a network protein of interest, is monitored by filter retardation ¹⁴. HD169Q68 *per se* has only a low propensity to form insoluble aggregates in HEK293 cells. However, as shown in Fig. 4a coexpression of the htt-interacting protein GIT1

strongly promotes the formation of HD169Q68 aggregates, whereas coexpression of PIASy, HIP5, HP28, PFN2, FEZ1 and BARD1 has no discernable effect. Thus, GIT1 is a potential modifier of HD pathogenesis, which may influence the rate of formation of insoluble htt aggregates *in vivo*.

Furthermore, probing of the insoluble HD169Q68 aggregates with an anti-GIT1 antibody revealed that GIT1 does not only stimulate aggregation but is also an integral part of the insoluble aggregates (Fig. 4a). This suggests that GIT1 promotes aggregation through direct binding to mutant htt.

The interaction between GIT1 and htt was confirmed by coimmunoprecipitation from COS1 cells transfected with constructs encoding HD510Q68 and HA-GIT1. Forty hours post transfection cell extracts were prepared and treated with antiserum against GIT1. HD510Q68 and HA-GIT1 were detected in the immunoprecipitate on Western blots with anti-htt antibody 4C8 and anti-HA antibody 12CA5, respectively (Fig. 4b).

The GIT1-htt interaction was also detected in human brain. Protein extracts prepared from human cortex were treated with the anti-htt antibodies CAG53b and HD1, and the precipitate was probed for the presence of GIT1 (Fig. 4c). Full length GIT1, migrating at about 90 kDa ³³, was precipitated by both anti-htt antibodies in a concentration dependent manner, indicating the existence of a complex between htt and GIT1 in neurons.

Finally, we performed colocalisation studies of htt and GIT1 in COS1 cells using immunofluorescence microscopy. In cells expressing HD510Q68 or GIT1 alone a diffuse cytoplasmic staining was observed for each protein (Fig. 4d). However, when GIT1 and mutant htt were coexpressed, large perinuclear structures, most likely reflecting protein aggregates, appeared almost exclusively. These structures contained both GIT1 and htt. The images further substantiate the findings that GIT1 and htt bind to each other and that GIT1 is a potent enhancer of mutant htt aggregation.

Example 5: GIT1 localises to htt aggregates in HD transgenic mouse and patient brains

The finding of colocalisation of htt and GIT1 within aggregates in transfected COS1 cells suggests that GIT1 might also be a component of htt aggregates in vivo. To investigate this possibility we first assessed the distribution of GIT1 in brains of R6/1 transgenic mice expressing a human htt exon 1 protein with 150 glutamines 34. In wildtype mice, GIT1 immunoreaction product was found diffuse in the cytoplasm and nuclei of neurons throughout the brain. In R6/1 brains, in addition to the diffuse staining, GIT1 immunoreactivity was also present in large nuclear and cytoplasmic puncta similar to htt aggregates (Figure 5a). To further confirm these data, we examined the subcellular distribution of GIT1 in cortex from HD patient brains and healthy individuals (Fig. 5b). In patient brains, GIT1 antibodies labelled neuronal nuclear inclusions as well as neuropil aggregates characteristic of HD brains 35. In contrast, neurons from control brains only showed a diffuse nuclear and cytoplasmic GIT1 immunostaining. In fact, in colocalisation studies performed in HD brain sections, GIT1 positive aggregates were also labelled with anti-htt antibody 2B4, indicating that both proteins coaggregated in vivo (Fig. 5c). This observation raises the possibility that an alteration of the neuronal GIT1 subcellular distribution contributes to HD pathogenesis.

PART II: VERIFICATION AND FURTHER RESULTS

Example 6: Experimental Procedures

Antibodies

A polyclonal antibody (pAb) against GIT1 was generated by injection of purified His6-tagged GIT1 (aa 368-587) into a rabbit. The resulting GIT1 pAb (C-GIT1) was affinity purified using immobilized GIT1 protein. The pAb NT-GIT1 recognizes the first 100 aa of GIT1 (Santa Cruz Biotechnology), the monoclonal antibody (mAb) CT-GIT1 (Transduction Laboratories) is specific for the last 10⁶ amino acids of GIT1. For all three Abs, no cross-reaction with GIT2 was observed (Fig. 13). The pAbs against GAPDH (Wanker et al., 1997) and htt [CAG53b (Davies et al., 1997) and HD1 (Scherzinger et al., 1997)] were described. Commercially available antibodies were anti-GST pAb (Amersham Pharmacia), anti-HA mAb 12CA5 (Roche Diagnostics), anti-htt pAb EM48 (Gutekunst et al., 1999), anti-htt mAb 2B4 (Lunkes et al., 2002), anti-htt mAb 4C8 (Chemicon) and anti-EEA1 pAb (Santa Cruz Biotechnology). As

secondary antibodies for immunofluorescence microscopy, Cy3- (dianova) and Alexa 488- (MoBiTec) conjugated IgGs were used.

Strains and plasmids

The yeast strains used for two-hybrid analysis were L40ccua [MATa his3D200 trp1-901 leu2-3,112 LYS2::(lexAop)4-HIS3 ura3::(lexAop)8-lacZ ADE2::(lexAop)8- URA3 GAL4 gal80 can1 cyh2] and L40cca [MATa his3D200 trp1-910 leu2-3,112 ade2 LYS2::(lexAop)4-HIS3 URA3::(lexAop)8-lacZ GAL4 gal80 can1 cyh2].

Plasmids pHD510Q17 and pHD510Q68 were generated by insertion of fragments coding for HD510Q17 and HD510Q68 into pcDNA-I (Invitrogen). pHD169Q68 was derived from pHD510Q68 by deletion of the Xhol-Xhol fragment encoding aa 170-510 of human htt. pV5-HD169Q68 was generated by inserting the EcoRI-Xhol fragment from pHD510Q68 into pcDNA3.1/V5-HIS (Invitrogen). Full-length GIT1 (aa 1-770) was amplified by PCR from the cDNA clone IMAGp958H111245Q2 (RZPD, Germany) using the primers GIT1-F/GIT1-R and subcloned into the EcoRI-BgIII site of pTL-HA (HA-GIT1). The GIT2 cDNA (aa 1-759) was PCR amplified with the primers GIT2-F / GIT2-R and subcloned into the Xhol-Notl site of pTL-HA (HA-GIT2). follows: GIT1-F The sequences were as primer GIT1-R (5'-GGAAGATCT CGGAATTCATGTCCCGAAAGGGGCCGCG-3'), GGTCACTGCTTCTCTCGGG-3'), GIT2-F (5'-ACGCGTCGACCATGTCGAAA CGGCTCCG-3') and GIT2-R (5'ATAAGAATGCGGCCGCGCCCTGCCCTTGCTA GTTG -3').

Library screening

Plasmids encoding baits were transformed into L40ccua, tested for the absence of reporter gene activity and cotransformed with a human fetal brain cDNA library (Clontech). For each transformation, 1 x 10⁶ independent transformants were plated onto minimal medium lacking tryptophan, leucine, histidine and uracil (SDIV medium) and incubated at 30°C for 5 to 10 days. Clones were picked into microtitre plates and

grown overnight in liquid minimal medium lacking tryptophan and leucine (SDII medium). Then, they were spotted onto nylon membranes placed on SDIV agar plates. After incubation for 4 days, the membranes were subjected to a b-galactosidase (b-GAL) assay. Plasmids were prepared from positive clones and characterized by sequencing. For retransformation assays, plasmids encoding baits and preys were cotransformed into L40ccua and plated onto SDIV medium.

• Array mating screen

Plasmids encoding baits and preys were transformed into strains L40ccua and L40cca, respectively. L40cca clones were arrayed in 96-well microtitre plates and mixed with a single L40ccua clone for interaction mating. Diploid cells were transferred onto YPD medium plates and, after incubation for 24 h at 30°C, onto SDII medium plates for additional 72 h at 30°C. For two-hybrid selection, diploid cells were transferred onto SDIV medium plates with and without nylon membranes and incubated for 5 days at 30°C. The nylon membranes were subjected to the b-GAL assay. Positive clones were verified by cotransformation assays.

Protein expression and verification assays

For verification experiments, cDNA fragments encoding baits and preys were subcloned into pGEX derivatives (Stratagene) or pTL-HA (Sittler et al., 1998). GST-fusion proteins were expressed in E. coli BL21-codon PlusTM RP (Stratagene) and affinity purified on glutathione agarose beads (Sigma) (Wanker et al., 1997). COS-1 cells were transfected with mammalian expression plasmids and lysed as described (Sittler et al., 1998). For in vitro binding assays, 30 µg of GST or GST fusion protein were immobilized on glutathione agarose beads and incubated with 500 µg COS-1 cell extract containing HA-tagged fusion protein for 2 h at 4°C, in binding buffer [50 mM HEPES-KOH pH 7.4, 150 mM NaCl, 10% glycerol, 1% NP-40, 1 mM EDTA, 20 mM NaF, 1 mM DTT, 0.1% Triton X-100, protease inhibitors (Roche Diagnostics)]. After centrifugation and extensive washing, bound proteins were eluted and analyzed by SDS-PAGE and Western blotting.

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Coimmunoprecipitation experiments were performed as previously described (Sittler et al., 1998). For immunofluorescence microscopy, COS-1 cells were grown on cover slips and cotransfected with plasmids encoding N-terminal htt V5-HD169Q68 and/or C-terminal HA-GIT1-CT. 40 h post-transfection, cells were treated with 2% paraformaldehyde. Immunolabeling was performed with anti-C-GIT1 (1:500) and with anti-V5 (1:300) Abs. Nuclei were counterstained with Hoechst. For subcellular localization of endogenous GIT1 and htt, differentiated PC12 and SH-SY5Y cells were used. PC12 cells were treated with 50 ng/ml NGF and grown on cover slips for 6 d. SH-SY5Y cells were serum starved for 24 h and then differentiated with 10 nM IGF-I for 30 min. Cells were labeled with C-GIT1 (1:20) and 4C8 (1:20) Abs and viewed with a confocal microscope LSM510 (Zeiss).

Filter Retardation Assay

HEK293 cells coexpressing HD169Q68 and selected network proteins were harvested 48 h post-transfection. Cell lysates were boiled in 2% SDS, 50 mM DTT for 5 min. Aliquots containing 12.5, 25 or 50 µg of total protein were used for filtration on cellulose acetate membranes (Scherzinger et al., 1997). SDS-resistant aggregates were detected using anti-CAG53b or anti C-GIT1 pAbs.

Inhibition of GIT1 expression by siRNA

For silencing of endogenous GIT1 expression, HEK293 cells were transfected with the siRNA duplex siRNA-GIT1 (5'-AAGCCTGGATGGAGACCTA GA-3') using TransMessenger (Qiagen) or Lipofectamin 2000 (Invitrogen) transfection reagents. 48 h post transfection, cell lysates were analyzed for GIT1 expression by immunoblotting using C-GIT1 Ab. To examine the effect of endogenous GIT1 silencing on htt aggregation, HEK293 cells were cotransfected with pHD169Q68 and siRNA-GIT1 and subjected to filtration after 72 h.

• Detection of GIT1 in R6/1 mouse and human HD brains ...

For immunocytochemistry, mice were deeply anaesthetized and perfused through the left cardiac ventricle with 4% paraformaldehyde in 0.1 M phosphate buffer. Brains were removed and postfixed overnight in 4% paraformaldehyde. Sections were processed for immunocytochemistry as described (Gutekunst et al., 1999). EM48 (1:1000) and C-GIT1 (1:100) pAbs were used.

Tissues from 8 human HD and 7 control brains were used in this study. Two HD cases were classified as grade 3 of neuropathological severity, six cases as grade 4. Standard protocols were used (Lunkes et al., 2002) for immunolabeling with 2B4 mAb (1:200) and C-GIT1 pAb (1:50). For Western analysis of total protein lysates from frontal cortex, the C-GIT1 pAb (1:300) was used.

Example 7:Two-hybrid screens

To generate a PPI network for HD, we used a combination of library and matrix yeast two-hybrid screens (Fig. 7A). Previous studies have shown that htt potentially participates in clathrin-mediated endocytosis, apoptosis, vesicle transport, cell signaling, morphogenesis and transcriptional regulation (Harjes and Wanker, 2003; Li and Li, 2004). For this reason, we selected 50 cDNAs encoding proteins involved in these processes, including 5 different N-terminal htt fragments, as well as proteins known to interact with htt, for subcloning into a DNA binding domain vector to express LexA fusion proteins as baits (Suppl. Table 1). The resulting plasmids were sequenced and introduced into yeast strain L40ccua, which carries three reporter genes, HIS3, URA3 and lacZ, for two-hybrid interaction analysis.

Forty of these baits did not activate the reporters by themselves and were used individually for cotransformation screening of a human fetal brain cDNA library expressing GAL4 activation domain (AD) hybrids as preys. In each screen, 1 x 10⁶ auxotrophic transformants were tested on selective plates, and 1-50 positive colonies were typically obtained. Restriction analysis and sequencing revealed that about 12% of all positive clones expressed preys with correct in-frame sequences, while 88% of the clones contained plasmids with out-of-frame sequences or sequences from non-protein-encoding regions, which were discarded. 27 preys were identified only once, while the other 11 were found up to four times as independent AD fusions. Plasmids

with the longest coding regions were used for subsequent studies. The preys identified by the library two-hybrid screens were tested together with their respective baits for activation of reporter gene expression in cotransformation assays. Only prey/bait combinations that activated the reporter gene expression in two independent cotransformation assays were selected for further two-hybrid studies and in vitro pull-down assays (Fig. 9). Starting with 40 baits in the library and subsequent cotransformation screens, we identified 41 PPIs among 18 bait and 38 prey proteins.

For a second round of two-hybrid screens, cDNAs encoding 12 prey proteins were selected from literature verified interactions and from interactions confirmed by in vitro binding experiments (Suppl. Table 2), and subcloned into a DNA binding domain vector. The resulting baits were tested for autoactivation, and 10 were screened against a human fetal brain cDNA library as described above. We identified another 14 PPIs among 5 bait and 13 prey proteins. Nine preys were found once and 4 were discovered multiple times as independent AD fusions. All interactions were confirmed by cotransformation assays.

Finally, an array-mating screen was performed to connect bait and prey proteins identified in the cDNA library transformation screens (Fig. 7A). L40cca yeast cultures were transformed with plasmids encoding the 51 prey proteins obtained in the first and second round of cDNA library screens and arrayed in 96-well microtiter plates. Prey cDNAs were also subcloned into DNA binding domain vectors and introduced into an L40ccua strain to generate additional baits for interaction mating. Including the ones already used for the library screens, we arrived at 46 baits, which did not autoactivate the reporter genes (Table 7). These baits were used individually for mating against the matrix of prey proteins. Diploid yeast clones, formed on YPD plates, were selected on agar SDII plates, and further transferred by a spotting robot onto SDIV plates to select for Y2H interactions (Fig. 7B). We examined 2346 (51x46) pair wise combinations of baits and preys in the mating assay reproducing all 55 two-hybrid interactions, which had been found in the library screens. In addition, 131 new PPIs were found by interaction mating and subsequently reproduced in cotransformation assays. Using this combination of library and matrix two-hybrid

screens, a total of 186 PPIs among 35 bait and 51 prey proteins could be identified (Fig. 8A).

Sequence analysis of the cDNAs revealed ORFs ranging from 82 to 728 amino acids in size (Table 7). In a systematic Blast search, 77 of the 86 bait and prey protein fragments were identical to a SwissProt or TrEMBL protein entry (http://us.expasy.org/sprot/). Nine proteins showed 75-99% identity to their best respective database hit and either contained single amino acid substitutions, variable polyQ lengths or small regions of sequence variation. Uncharacterized proteins were named according to their interaction partners.

This chapter describes the whole yeast two hybrid screening procedure and obtained fundamental data. A full description of our final datasets are shown in tables 6 to 9. Table 6 contains a compilation of all found protein—protein interactions in the Huntington's disease protein network. Some of these interactions are already known and literature-cited. A dataset which describes only new identified interactions will be found in Table 9. Table 7 characterizes all proteins involved in the protein network. Most of these proteins are known from different databases but some proteins are still unknown (Table 8). Nucleic acid and amino acid sequence data for all network-proteins are available from Figure 6.

Example 8: Functional assignment of yeast two-hybrid data

To chart two-hybrid interactions identified in this study, previously reported, or verified by independent methods, a matrix representation was used (Fig. 8A). We assigned a subcellular localization to each network protein by examining various sources of literature and, based on the experimental data, we grouped the proteins into six broad functional categories (Fig. 8A, Table 7). 18 proteins in the HD network are involved in transcriptional regulation or DNA maintenance; 14 proteins mainly participate in cytoskeletal and transport processes. We assigned 7 proteins to cellular signaling and fate, another 5 to protein synthesis and turnover, and 3 proteins to cellular metabolism. 16 proteins of unknown function were identified, participating in 72 interactions. The number of interactions identified for each protein varied from 1 to 24, with 2.6 interaction partners on average. Interestingly, proteins such as htt,

BARD1, GADD45G, HIP5, HZFH, PIASy, BAIP3 or VIM interact with more than 15 other proteins, forming hubs in the HD network, while 15 proteins have only one interaction partner.

For htt, 19 different interacting partners from various functional groups were identified, of which HIP1, CA150, SH3GL3 and HYPA had been described previously (Harjes and Wanker, 2003). 6 of the htt partners are involved in transcriptional regulation and DNA maintenance, 4 function in cellular organization and transport and 3 in cellular signaling, supporting the hypothesis that htt is involved in these processes. Moreover, we have detected 6 novel htt interacting proteins of unknown function termed HIP5, HIP11, HIP13, HIP15, HIP16, and CGI-125.

Using 5 different N-terminal htt fragments as baits, the potential htt-binding sites of 13 interaction partners were mapped (Fig. 8A). For the proteins CA150, HYPA, PNF2, SH3GL3, CGI-125 and HIP13, however, a conclusive determination of the htt binding region was not possible with the two-hybrid assay, because these proteins bound to HDexQ20, HDexQ51 and HD1.7, but not to HDd1.0 (Fig. 8A). We suggest that these proteins bind to the htt exon 1 fragment, but this binding region might be masked in the HDd1.0 protein, while it is accessible in the HD1.7 fragment. Interestingly, we found that HP28 and HIP15 bind to HDexQ51, but not to HDexQ20, HD1.7 and HD1.0, indicating that the interaction of these proteins with htt is enhanced by the expanded polyQ repeat. Thus, HP28 and HIP15 may be disease specific htt interactors.

To generate a more comprehensive HD interaction map, we supplemented our two-hybrid network (red diamonds) with all 38 known direct htt interaction partners (Suppl. Table 4 and Fig. 8B, blue squares). Furthermore, we added 83 human proteins (green triangles), identified from protein interaction databases HPRD, MINT, and BIND that bridge any two proteins in our extended network. Using this approach, we obtained an interaction network for htt containing a total of 181 proteins and 591 PPIs (Fig. 8B and Suppl. Table 5).

Example 9: Verification of PPIs

Comparison with literature-cited interactions revealed that more than 89% of the twohybrid interactions identified are unknown. 30 PPIs have been reported previously using two-hybrid methods, coimmunoprecipitations or affinity chromatography-based techniques; 21 of these were detected in our Y2H assays (Fig. 8A, Suppl. Table 3). In most cases, the occurrence of false negatives can be explained by the lack of essential domains in one of the protein fragments. For example, an interaction between p53 and hADA3 has been described (Wang et al., 2001), with the first 214 amino acids of hADA3 being essential for this interaction. It escaped our two-hybrid analysis, because a C-terminal hADA3 fragment (amino acids 235-432) was used.

Failure to detect interactions may also result from the high stringency of our twohybrid assay, which can be attributed to low protein expression levels and the simultaneous use of three reporters. Our system is particularly designed to minimize false positives, which are known to occur frequently in two-hybrid assays (von Mering et al., 2002). To determine the rate of false positives in our system, we directly assessed 54 interactions from the two-hybrid network by in vitro pull-down experiments, mainly focusing on htt and its immediate interaction partners. Proteins were expressed as GST-fusions in E. coli, and their interacting partners as HAfusions in COS-1 cells. After immobilization of GST-fusion proteins to beads, the potential interaction partners were pulled down from COS-1 cell extracts. Binding was detected by SDS-PAGE and immunoblotting. Using this assay, 35 interactions representing 32 different protein pairs were verified successfully (Fig. 9). Failure to detect an interaction by GST pull-down assays could be due to low protein expression levels or the lack of appropriate protein modifications. Therefore, the 19 non-verified protein-protein interactions are still valid until further experiments show contradictory results. The rate of 64.8% verified interactions suggests that in our Y2H network false positives might appear less frequently than described for other PPI studies (von Mering et al., 2002).

Example 10: GIT1 promotes htt aggregation

Several lines of evidence indicate that aggregation of mutant htt is linked to disease progression and the development of motor symptoms (Davies et al., 1997; Sanchez et al., 2003). Therefore, cellular proteins that influence aggregate formation are potential modulators of disease pathogenesis. In order to identify such proteins, we screened all 19 direct htt interaction partners (Fig. 8A) for their ability to enhance htt aggregation in a cell-based assay (Sittler et al., 1998). In this assay, HEK293 cells were cotransfected with constructs encoding an aggregation prone N-terminal htt fragment with 68 glutamines (HD169Q68) and a network protein. After 48 h, formation of SDS-insoluble htt aggregates was analyzed by a filter retardation assay (Scherzinger et al., 1997). In this time period HD169Q68 by itself formed only few aggregates. In comparison, coexpression of the C-terminal GIT1 fragment found in the Y2H screens (GIT1-CT) increased the amount of htt aggregates 3-fold (Fig. 10A). Coexpression of HD169Q68 with other htt-interacting proteins, on the other hand, did not enhance htt aggregation (data not shown).

It has been described previously that GIT1 and its homologue p95-APP1 are able to form homo- and heterodimers in vitro and in vivo (Kim et al., 2003; Paris et al., 2003). Therefore, we wondered whether GIT1-CT by itself is able to form SDS-insoluble protein aggregates in mammalian cells. As shown in Fig. 10A, we did not detect aggregates in the filter retardation assay upon transient overexpression of GIT1-CT. However, in cells coexpressing HD169Q68 and GIT1-CT, stable SDS-resistant aggregates immunoreactive with the anti-C-GIT1 antibody were formed, indicating that both proteins coaggregate in cells, and that GIT1-CT is an integral part of the insoluble htt aggregates (Fig. 10A).

Next, we tested whether full-length GIT1 is able to accelerate htt aggregation in mammalian cells. Analysis by filter retardation assay revealed that full-length GIT1 enhances htt aggregation in a dose dependent manner (Fig. 10B). However, compared to GIT1-CT, it was less efficient in stimulating HD169Q68 aggregation in the cell model, indicating that the N-terminally truncated GIT1 fragment is a more potent enhancer of htt aggregation than the full-length protein.

As previous studies have shown that the expression of C-terminal GIT1/p95-APP1 fragments induces the formation of large vesicular structures in mammalian cells (Di Cesare et al., 2000; Matafora et al., 2001), we analyzed the effect of GIT1-CT on HD169Q68 aggregation by indirect immunofluorescence microscopy. We found that expression of GIT1-CT alone induced the accumulation of large vesicular structures

in the perinuclear region (Fig. 10Cb). In comparison, when HD169Q68 was expressed alone, the protein was distributed in the cytoplasm, and no large aggregates or inclusion bodies were observed (Fig. 10Ca). However, when HD169Q68 and GIT1-CT were coexpressed (Fig. 10Cd-f), htt was almost exclusively detected in the perinuclear vesicles (Fig. 10Cd), indicating that GIT1-CT overexpression induces the relocalization of htt to membranous structures. A similar effect was observed when full-length GIT1 and HD169Q68 were coexpressed in COS1 cells, however, the rate of vesicle formation and htt recruitment was lower, compared to GIT1-CT/HD169Q68 expressing cells (data not shown). The colocalization of GIT1 with the early endosomal marker EEA1 is shown in Fig. 10Cc. Together, these results suggest that the enhancement of HD169Q68 aggregation in mammalian cells is due to the recruitment of mutant htt into vesicular structures induced by overexpression of GIT1 or GIT1-CT.

Example 11: GIT1 is crucial for the formation of htt aggregates in mammalian cells

Next, we investigated whether endogenous GIT1 promotes htt aggregation in mammalian cells. In order to reduce endogenous GIT1 levels in HEK293 cells, we employed the short-interfering RNA (siRNA) technology (Elbashir et al., 2001). Cells were cotransfected with HD169Q68 and GIT1-specific siRNA, and silencing of endogenous GIT1 was monitored 48 h post transfection by Western blot analysis (Fig. 10D). We found that siRNA treatment specifically reduced endogenous GIT1 by ~80% and caused a strong decrease of HD169Q68 aggregate formation (Fig. 10E). After incubation for 72 h, SDS-resistant HD169Q68 aggregates were detected in untreated, but not in siRNA treated cells. This indicates that physiological levels of GIT1 are critical for htt aggregation in mammalian cells, and that an inhibition of GIT1 expression dramatically slows down aggregate formation. A similar effect was also obtained when GIT1-specific siRNA was applied to cells overexpressing GIT1-CT and HD169Q68 proteins (data not shown).

Example 12: Verification of the htt-GIT1 interaction

The interaction between GIT1-CT and htt was confirmed by coimmunoprecipitation from COS-1 cells transfected with constructs encoding the first 510 amino acids of htt with 68 glutamines (HD510Q68) and an N-terminally truncated hemagglutinin (HA) tagged HA-GIT1-CT (aa 249-770) protein. 40 h post-transfection, cell extracts were prepared and treated with GIT1 antiserum. HD510Q68 and HA-GIT1-CT were detected in the immunoprecipitates on Western blots with anti-htt antibody 4C8 and anti-HA antibody 12CA5, respectively (Fig. 11A).

The GIT1-htt interaction was also detected in healthy human brain. Protein extracts prepared from cortex were treated with the anti-htt antibodies CAG53b and HD1, and the precipitate was probed for the presence of GIT1 (Fig. 11B) with a GIT1 specific antibody (NT-GIT1; Fig. 13). Full length GIT1, migrating at about 95 kDa (Vitale et al., 2000), was precipitated by both anti-htt antibodies in a concentration dependent manner, indicating that a protein complex containing htt and GIT1 is formed under physiological conditions.

Next, we examined the colocalization of endogenous htt and GIT1 in differentiated PC12 cells by confocal immunofluorescence microscopy. Both proteins were mainly detected in the cytoplasm, but were also present in the neurite-like extensions (Fig. 11Cab). Colocalization, indicated in yellow, was visible in cytoplasmic complexes in the perinuclear region (Fig. 11Cc) as well as in a number of intracellular structures scattered throughout the neuritic extensions. GIT1 was also detected in adhesion-like structures at the tip of the extensions, as previously reported (Di Cesare et al., 2000; Manabe Ri et al., 2002). These regions, however, did not contain htt protein. Similar results were obtained when the endogenous localization of GIT1 and htt was analyzed in differentiated neuroblastoma SH-SY5Y cells using confocal immunofluorescence microscopy (Fig. 11Cd-f).

Example 13: GIT1 localizes to htt aggregates in patient brain

Our findings suggest that GIT1 might also be a component of neuronal inclusions containing htt aggregates in brain of HD patients and transgenic animals (Davies et al., 1997; DiFiglia et al., 1997). To investigate this possibility, we first assessed the distribution of GIT1 in brain slices of R6/1 transgenic mice expressing a human htt exon 1 protein with 150 glutamines (Mangiarini et al., 1996). In wild type mice, GIT1 specific immunoreactivity was diffused in the cytoplasm and nuclei of neurons throughout the brain. In R6/1 brain, however, in addition to a diffuse staining, GIT1 immunoreactivity was also present in large nuclear and cytoplasmic puncta containing htt aggregates (Fig. 12A). To further confirm these data, we examined the subcellular distribution of GIT1 in HD patient and healthy cortex (Fig. 12B). In patient brain, GIT1 specific antibodies labeled neuronal nuclear inclusions as well as the neuropil aggregates characteristic of HD (DiFiglia et al., 1997). In contrast, neurons from control tissue showed only diffuse nuclear and cytoplasmic GIT1 immunostaining. Fig. 12C shows colocalization of htt and GIT1 in neuronal nuclear

Example 14: GIT1 is degraded in HD patient brain

inclusions.

The presence of GIT1 in protein extracts from HD affected and unaffected cortex was also analyzed by SDS-PAGE and immunoblotting. As shown in Fig. 12D, full-length GIT1 protein migrating at about 95 kDa was detected in healthy brain (Fig. 12D), but was significantly reduced in HD. Interestingly, in HD, but not in control brain, prominent GIT1 degradation products migrating at about 25-50 kDa were detected with the C-terminal GIT1 antibody C-GIT1 (Fig. 12D). In strong contrast, no such products were observed when the N-terminal GIT1 antibody NT-GIT1 directed against the ARF-GAP domain was used (data not shown). This indicates the formation of large amounts of N-terminally truncated GIT1 fragments in HD brain, which may be a significant factor in disease pathogenesis.

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Claims

- 1. A method for generating a network of direct and indirect interaction partners of a disease-related (poly)peptide comprising the steps of
 - (a) contacting a selection of (poly)peptides suspected to contain one or several of said direct or indirect interaction partners with said disease-related (poly)peptides and optionally with known direct or indirect interaction partners of said disease-related (poly)peptide under conditions that allow the interaction between interaction partners to occur;
 - (b) detecting (poly)peptides that interact with said disease-related (poly)peptide or with said known direct or indirect interaction partners of said disease-related (poly)peptide;
 - (c) contacting (poly)peptides detected in step (b) with a selection of (poly)peptides suspected to contain one or several (poly)peptides interacting with said (poly)peptides detected in step (b) under conditions that allow the interaction between interaction partners to occur;
 - (d) detecting proteins that interact with said (poly)peptides detected in step (b);
 - (e) contacting said disease-related (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide, said (poly)peptides detected in steps (b) and (d) and a selection of proteins suspected to contain one or several (poly)peptides interacting with any of the afore mentioned (poly)peptides under conditions that allow the interaction between interaction partners to occur;
 - (f) detecting (poly)peptides that interact with said disease-related (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide or with said (poly)peptides identified in step (b) or (d); and
 - (g) generating a (poly)peptide-(poly)peptide interaction network of said disease-related (poly)peptide and optionally said known direct or indirect interaction partners of said disease-related (poly)peptide and said (poly)peptides identified in steps (b), (d) and (f).
- 2. The method of claim 1, wherein said contacting step (e) is effected in an interaction mating two hybrid approach.

- 3. The method of claim 1 or 2, said method comprising after step (d) and before step (e) the steps of:
 - (d') contacting (poly)peptides detected in step (d) with a selection of (poly)peptides suspected to contain one or several (poly)peptides interacting with said (poly)peptides detected in step (d) under conditions that allow the interaction between interaction partners to occur; and
 - (d") detecting proteins that interact with said (poly)peptides detected in step (d').
- 4. The method of any one of claims 1 to 3, wherein said disease-related protein is a protein suspected of being a causative agent of a hereditary disease.
- 5. The method of any one of claims 1 to 4, wherein said disease-related protein is huntingtin and wherein said interaction partners are the interaction partners as shown in tables 6, 7 or 9.
- 6. The method of any one of claims 1 to 5, said method comprising the step of determining the nucleotide sequence of a nucleic acid molecule encoding a direct or indirect interaction partner of the disease related protein.
- 7. The method of any one of claims 1 to 6, wherein said selections of proteins are translated from a nucleic acid library.
- 8. The method of any one of claims 1 to 7, wherein said selection of proteins in step (a) and/or (c) and/or (d') and/or (e) is the same selection or a selection from the same source.
- 9. The method of any one of claims 1 to 7, wherein said selection of proteins in step (a) and/or (c) and/or (d') and/or (e) is a different selection or a selection from a different source.
- 10. The method of any one of claims 1 to 9, wherein said method is performed by contacting the proteins on an array.

- 11. The method of any one of claims 1 to 10, wherein said interactions are detected by using the yeast two-hybrid system.
- 12. The method of any one of claims 1 to 11, containing after step (b), (d), (d") or (f) the additional steps of isolating a nucleic acid molecule with homology to said cDNA expressing the encoded protein and testing it for its activity as a modulator of huntingtin, wherein said nucleic acid molecule is DNA, or RNA, preferably cDNA, or genomic or synthetic DNA or mRNA.
- 13.A nucleic acid molecule encoding a modulator of huntingtin, wherein said modulator is a protein selected from table 8.
- 14. The nucleic acid molecule of claim 13, wherein said nucleic acid molecule is DNA, preferably cDNA, genomic DNA, or synthetic DNA or RNA, preferably mRNA.
- 15. The nucleic acid molecule of claim 13 or 14 fused to a heterologous nucleic acid molecule.
- 16. The nucleic acid molecule of claim 15, wherein the heterologous nucleic acid molecule encodes a heterologous (poly) peptide.
- 17.A vector comprising the nucleic acid molecule of any one of claims 13 to 16.
- 18.A host cell containing the nucleic acid molecule of any one of claims 13 to 16 or the vector of claim 17.
- 19.A method of producing a (poly)peptide, comprising culturing the host cell of claim 18 under conditions such that the (poly)peptide encoded by said polynucleotide is expressed and recovering said (poly)peptide.
- 20.A (poly)peptide comprising an amino acid sequence encoded by a nucleic acid molecule of any one of claims 13 to 16, or which is chemically synthesized, or is obtainable from the host cell of claim 18, or which is obtainable by the method of

claim 19 or which is obtainable from an in vitro translation system by expressing the nucleic acid molecule of any one of claims 13 to 16 or the vector of claim 17.

- 21. The (poly)peptide of claim 20 fused to a heterologous (poly)peptide.
- 22.A protein complex comprising at least two proteins, wherein said at least two proteins are selected from the group of interaction partners listed in table 9.
- 23.An antibody specifically recognizing the (poly)peptide of claim 20 or 21 or specifically reacting with the protein complex of claim 22.
- 24. The antibody of claim 23 which is polyclonal, monoclonal, chimeric, single chain, single chain Fv, human antibody, humanized antibody, or Fab fragment.
- 25.A method of identifying whether a protein promotes huntingtin aggregation, comprising
 - transfecting a first cell with a nucleic acid molecule encoding a variant of the huntingtin protein or a fragment thereof capable of forming huntingtin aggregates;
 - (b) co-transfecting a second cell with
 - (i.) a nucleic acid molecule encoding a variant of the huntingtin protein or a fragment thereof capable of forming huntingtin aggregates; and
 - (ii.) a nucleic acid molecule encoding a candidate modulator protein identified by the methods of any one of claims 1 to 12 or a nucleic acid molecule encoding a modulator protein selected from table 6 or table 7:
 - (c) expressing the proteins encoded by the transfected nucleic acid molecule of (a) and (b);
 - (d) isolating insoluble aggregates of huntingtin from the transfected cell of (a) and (b); and
 - (e) determining the amount of insoluble huntingtin aggregates from the transfected cell of (a) and (b)
 - wherein an increased amount of huntingtin aggregates isolated from the transfected cells of (b) in comparison with the amount of huntingtin aggregates

isolated from the transfected cells of (a) is indicative of a protein's activity as an enhancer of huntingtin aggregation.

- 26.A method of identifying whether a protein inhibits huntingtin aggregation, comprising
 - (a) transfecting a first cell with a nucleic acid molecule encoding a variant of the huntingtin protein or a fragment thereof capable of forming huntingtin aggregates;
 - (b) co-transfecting a second cell with
 - (i.) a nucleic acid molecule encoding a variant of the huntingtin protein or a fragment thereof capable of forming huntingtin aggregates; and
 - (ii.) a nucleic acid molecule encoding a candidate modulator protein identified by the methods of any one of claims 1 to 12 or a nucleic acid molecule encoding a modulator protein selected from table 6 or table 7;
 - (c) expressing the proteins encoded by the transfected nucleic acid molecule of (a) and (b);
 - (d) isolating insoluble aggregates of huntingtin from the transfected cell of (a) and (b); and
 - (e) determining the amount of insoluble huntingtin aggregates from the transfected cell of (a) and (b)

wherein a reduced amount of huntingtin aggregates isolated from the transfected cells of (b) in comparison with the amount of huntingtin aggregates isolated from the transfected cells of (a) is indicative of a protein's activity as an inhibitor of huntingtin aggregation.

- 27. The method of claim 25 or 26, wherein prior to step (d) the cells are treated with an ionic detergent.
- 28. The method of any one of claims 25 to 27, wherein the huntingtin aggregates are filtered or transferred onto a membrane.

- 29. A method for identifying compounds affecting an interaction of huntingtin or of a direct or indirect interaction partner of huntingtin comprising
 - (a) contacting interacting proteins selected from the group of interacting proteins listed in table 6 and/or table 7 in the presence or absence of an potential modular of interaction;
 - (b) identifying compounds capable of modulating said interaction.
- 30. The method of any one of claims 25 to 29, further comprising
 - (a) modeling said compound by peptidomentics and
 - (b) chemically synthesizing the modeled compound.
- 31. The method of any one of claims 25 to 30, wherein said compound is further modified to achieve
 - (i) modified site of action, spectrum of activity, organ specificity, and/or
 - (ii) improved potency, and/or
 - (iii) decreased toxicity (improved therapeutic index), and/or
 - (iv) decreased side effects, and/or
 - (v) modified onset of therapeutic action, duration of effect, and/or
 - (vi) modified pharmakinetic parameters (resorption, distribution, metabolism and excretion), and/or
 - (vii) modified physico-chemical parameters (solubility, hygroscopicity, color, taste, odor, stability, state), and/or
 - (viii) improved general specificity, organ/tissue specificity, and/or
 - (ix) optimized application form and route

p.

- (i) esterification of carboxyl groups, or
- (ii) esterification of hydroxyl groups with carbon acids, or
- (iii) esterification of hydroxyl groups to, e.g. phosphates, pyrophosphates or sulfates or hemi succinates, or
- (iv) formation of pharmaceutically acceptable salts, or
- (v) formation of pharmaceutically acceptable complexes, or
- (vi) synthesis of pharmacologically active polymers, or
- (vii) introduction of hydrophilic moieties, or

- (viii) introduction/exchange of substituents on aromates or side chains, change of substituent pattern, or
- (ix) modification by introduction of isosteric or bioisosteric moieties, or
- (x) synthesis of homologous compounds, or
- (xi) introduction of branched side chains, or
- (xii) conversion of alkyl substituents to cyclic analogues, or
- (xiii) derivatisation of hydroxyl group to ketales, acetales, or
- (xiv) N-acetylation to amides, phenylcarbamates, or
- (xv) synthesis of Mannich bases, imines, or
- (xvi) transformation of ketones or aldehydes to Schiff's bases, oximes, acetales, ketales, enolesters, oxazolidines, thiozolidines or combinations thereof.
- 32. A method of diagnosing Huntington's disease in a biological sample comprising the steps of
 - (a) contacting the sample with an antibody specific for a protein of table 6or 7 or an antibody specific for the protein complex of claim 22; and
 - (b) detecting binding of the antibody to a protein complex, wherein the detection of binding is indicative of Huntington's disease or of a predisposition to develop Huntington's disease.
- 33. The method of claim 32, wherein
 - (a) said protein complex contains GIT1 or
 - (b) said antibody is specific for a protein complex containing GIT1.
- 34. The method of claim 32, wherein
 - (a) said protein complex contains at least one protein selected from htt, HIP15 or HP28
 - (b) said antibody is specific for a protein complex containing at least one protein selected from htt, HIP15 or HP28.
- 35. A diagnostic agent/composition or pharmaceutical composition comprising the nucleic acid molecule of any one of claims 13 to 16, the (poly)peptide of claim 20 or 21 or the (poly)peptide mentioned in anyone of tables 6 and 7, the

- antibody of claim 23 or 24, an antibody specifically reacting with a protein selected from table 7 and/or a protein selected from table 7.
- 36. Use of the molecule of any one of claims 13 to 16, the (poly)peptide of claim 20 or 21 or the (poly)peptide mentioned in anyone of tables 6 and 7, the antibody of claim 23 or 24, an antibody specifically reacting with a protein selected from table 7 and/or a protein selected from table 7, for the preparation of a pharmaceutical composition for the treatment of Huntington's disease.

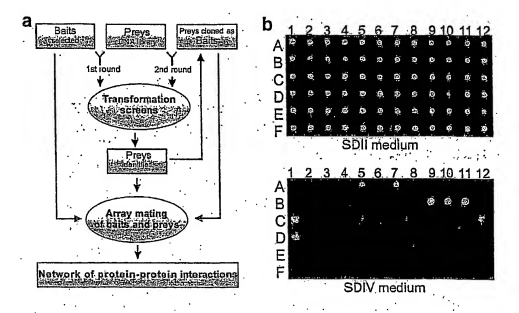
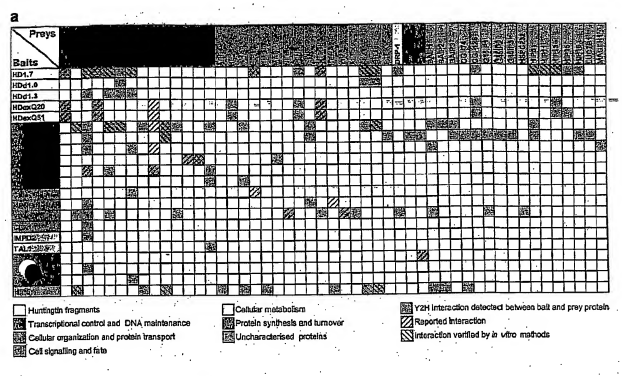


Figure 1



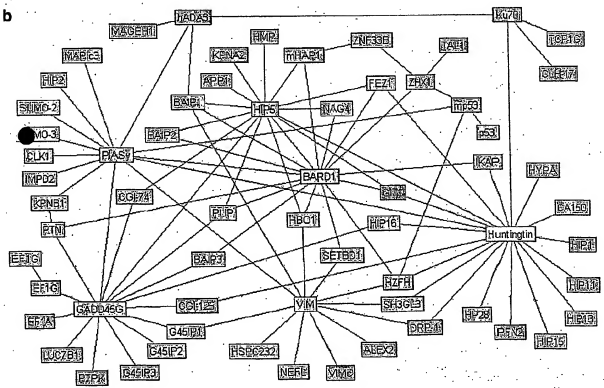
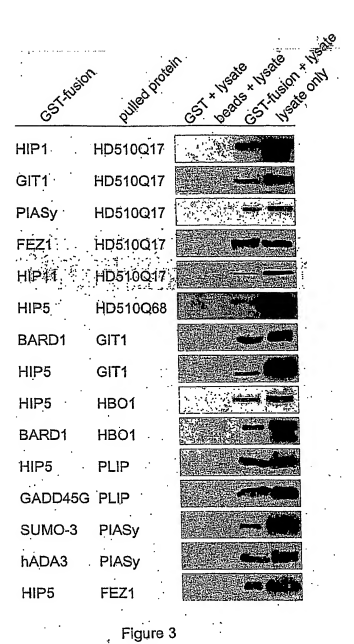
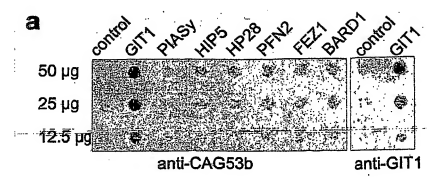
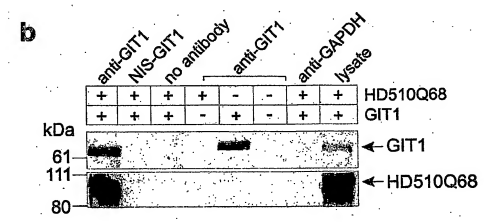
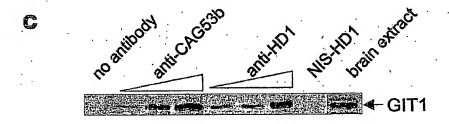


Figure 2









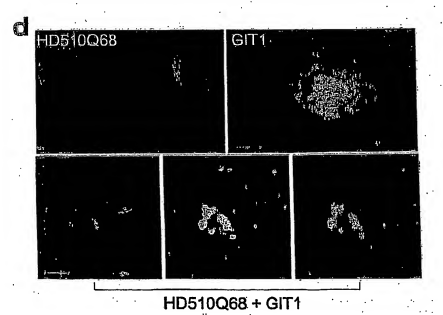


Figure 4

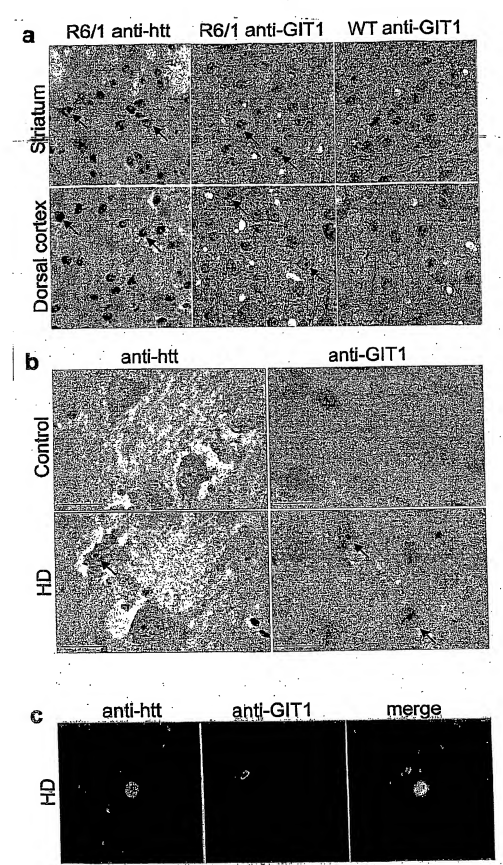


Figure 5

>ALEX2

AESVVGAAMASAIAPPPGVTEALGAAEAPAMAGAPKVAEAPREAETSRAAVPPGTVVPTEAZAPTE
VTEGPGVAAPTKVAEAPGVASPTEAAEAPVPATPTGAAAPTGAAESPGTSGSPRTAVVPGTSAAKK
ATPGAHTGAIPKATSATGAVPKGGGKGVTRSRNGGKGKGKKSKVEVDELGMGFRPGDGAAAZAAAS
ANGGQAFLAEVPDSEEGESGWTDTESDSDSEPETQRRGRGRRPVAMQKRPFPYEIDEILGVRDLRK
VLALLQKSDDPFIQQVALLTLSNNANYSCNOETIRKLGGLPIIANMINKTDPHIKEKALMAMMNLS
ENYENGGRLQVYMNKVMDDTMASNINSAVQVVGLKFLTNMTITNDYQHLLVNSIANFFRLLSQGGG
KIKVEILKILSNFAENPDMLKKLLSTQVPASFSSLYNSYVESEILINALTLFEIIYDNLRAEVFNY
REFNKGSLFYLCTTSGVCVKKIRALANHHDLLVKVKVIKLVNKF

EEEEESFPQPVDDYFVEPPQAEEEEETVPPPSSHTLAVVGKVTPTPRPTDGVDIYFGMPGEISEHE GFLRAKMDLEERRMRQINEVMREWAMADNQSKNLPKADRQALMEHFQSILQTLEEQVSGERQRLVE THATRVIALINDQRRAALEGFLAALQADPPQAERVLLALRRYLRAEQKEQRHTLRHYQHVAAVDPE KAQQMRFQVHTHLQVIEERVNQSLGLLDQNPHLAQELRPQIQELLHSEHLGPSELEAPAPGGSSED KGGLQPPDSKDDTPMTLPKGSTEQDAASPEKEKMNPLEQYERKVNASVPGVSLSTHRRFRGMSWHQLGQGCPVRLCRVC

>BAIP1

RPRTKMATAMYLEHYLDSIENLPCELQRNFQLMRELDQRTEDKKAEIDILAAEYISTVKTLSPDQR VERLQKIQNAYSKCKEYSDDKVQLAMQTYEMVDKHIRRLDADLARFEADLKDKMEGSDFESSGGRG LKKGRGQKEKRGSRGRGRRTSEEDTPKKKKHKGGSEFTDTILSVHPSDVLDMPVDPNEPTYCLCHQ VSYGEMIGCDNPDCPIEWFHFACVDLTTKPKGKW

>BAIP2

SQQASVTMHDVDAESFEVLVDYCYTGRVSLSEANVQRLYAASDMLQLEYVREACASFLARRLDLTN CTAILKFADAFDHHKLRSQAQSYLAHNFKQLSRMGSIREETLADLTLAQLLAVLRLDSLDIESERT VCHVAVQWLEAAAKERGPSAAEVFKCVRWMHFTEEDQDYLEGLLTKPIVKKYCLDVIEGALQMRYG DLLYKSLVPVPNSSSSSSSNSLVSAAENPPQRLGMCAKEMVIFFGHPRDPFLCYDPYSGDIYTMP SPLTSFAHTKTVTSSAVCVSPDHDIYLAAQPRKDLWVYKPAQNSWQQLADRLLCREGMDVAYLNGY IYILGGRDPITGVKLKEVECYSVQRNQWALVAPVPHSFYSFELIVVQNYLYAVNSKRMLCYDPSHN MWLNCASLKRSDFQEACVFNDEIYCICDIPVMKVYNPARGEWRRISNIPLDSETHNYQIVNHDQKL LLITSTTPQWKKNRVTVYEYDTREDQWINIGTMLGLLQFDSGFICLCARVYPSCLEPGQSFITEED DARSESSTEWDLDGFSELDSESGSSSFSDDEVWVQVAPQRNAQDQQGSL

>BAIP3

GHNAPRKVTAVIYARKGSVLQSIEKISSSVDATTVTSQQCVFRDQEPKIHNEMASTSDKGAQGRND KKDSQGRSNKALHLKSDAEFKKIFGLTKDLRVCLTRIPDHLTSGEGFDSFSSLVKSGTYKETEFMV KEGERKQQNFDKKRKAKTNKKMDHIKKRKTENAYNAIINGEANVTGSQLLSSILPTSDVSQHNILT SHSKTRQEKRTEMEYYTHEKQEKGTLNSNAAYEQSHFFNKNYTEDIFPVTPPELEETIRDEKIRRL KQVLREKEAALEEMRKKMHQK

~BARD1

LAGFESLTCSFPVVSRGLLASRSPRSLSSEGGTMPDNRQPRNRQPRIRSGNEPRSASAMEPDGRGA WAHSRAALDRLEKLLRCSRCTNILREPVCLGGCEHIFCSNCVSDCIGTGCPVCYTPAWIQDLKINR QLDSMIQLCSKLRNLLHDNEPSDLKEDKPRKSLFNDAGNKKNSIKMWFSPRSKKVRYVVSKASVQT QPAIKKDASAQQDSYEFVSPSPPADVSERAKKASARSGKKQKKKTLAEINQKWNLEAEKEDGEFDS KEESKQKLVSFCSQPSVISSPQINGEIDLLASGSLTESECFGSLTEVSLPLAEQIESPDTKSRNEV VTPEKVCKNYLTSKKSLPLENNGKRGHHNRLSSPISKRCRTSILSTSGDFVKQTVPSENIPLPECS SPPSCKRKVGGTSGSKTVTCPMNSLVFHQVHHLLH

>CA150

QQFIPGPLKILVWPCCLFSQAPTTQDQTPSSAVSVATPTVSVSTPAPTATPVQTVPQPHPQTLPPA VPHSVPQPTTAIPAFPPVMVPPFRVPLPGMPIPLFGVLPGMAPPIVPMIHPQVAIAASPATLAGAT AVSEWTEYKTADGKTYYYNNRTLESTWEKPQELKEKEKLEEKIKEPIKEPSEEPLPMETEEEDPKE EPIKEIKEEPKEEEMTEEEKAAQKAKPVATAPIPGTPWCVVWTGDERVFFYNPTTRLSMWDRPDDL IGRADVDKIIQEPPHKKGMEELKKLRHPTPTMLSIQKWQFSMSAIKEEQELMEEINEDEPVKAKKR WO 2004/113566 PCT/EP2004/006617 7/51

>CGI-125

PDASARNFARVSGLLLCQAGGVLVSSFVMAAAVAMETDDAGNRLRFQLELEFVQCLANPNYINFLA QRGYFKDKAFVNYLKYLLYWKDPEYAKYLKYPQCLHMLELLQYEHFRKELVNAQCAKFIDEOQILH WOHYERKRMRLQQALAEQQQQNNTSGK

>CGI-74

<u>VEKARAKKREAEEVYRNSMPASSFOOOKLRVCEVCSAYLGLHDNDRRLADHFGGKLHLGFIEIREK.</u> **DEEDKRYVAEKQEKRNQERLKRREEREREREKLRRSRSHSKNPKR**

>CLH-17

MAQILPIRFQEHLQLQNLGINPANIGFSTLTMESDKFICIREKVGEOAOVVIIDMNDPSNPIRRPI SADSAIMNPASKVIALKAGKTLQIFNIEMKSKMKAHTMTDDVTFWKWISLNTVALVTDNAVYHWSM EGESQPVKMFDRHSSLAGCQIINYRTDAKQKWLLLTGISAQQNRVVGAMQLYSVDRKVSQPIEGHA ASFAQFKMEGNAEESTLFCFAVRGQAGGKLHIIEVGTPPTGNQPFPKKAVDVFFPPEAQNDFPVAM QISEKHDVVFLITKYGYIHLYDLET

>CLK1

DAWVLEHLNTTDPNSTFRCVOMLEWFEHHGHICIVFELLGLSTYDFIKENGFLPFRLDHIRKMAYO ICKSVNFLHSNKLTHTDLKPENILFVQSDYTEAYNPKIKRDERTLINPDIKVVDFGSATYDDEHHS TLVSTRHYRAPEVILALGWSQPCDVWSIGCILIEYYLGFTVFPTHDSKEHLAMMERILGPLPKHMI QKTRKRKYFHHDRLDWDEHSSAGRYVSRRCKPLKEFMLSQDVEHERLFDLIQKMLEYDPAKRITLR EALKHPFFDLLKKSI

>DRP-1

KDNFTLIPEGVNGIEERMTVVWDKAVATGKMDENQFVAVTSTNAAKIFNLYPRKGRIAVGSDADVV IWDPDKLKTITAKSHKSAVEYNIFEGMECHGSPLVVISQGKIVFEDGNINVNKGMGRFIPRKAFPE HLYQRVKIRNKVFGLQGVSRGMYDGPVYEVPATPKYATPAPSAKSSPSKHQPPPIRNLHQSNFSLS GAOIDDNNPRRTGHRIVAPPGGRSNITSLG

MHHEALSEALPGDNVGFNVKNVSVKDVRRGNVAGDSKNDPPMEAAGFTAQVIILNHPGQISAGYAP VLDCHTAHIACKFAELKEKIDRRSGKKLEDGPKFLKSGDAAIVDMVPGKPMCVESFSDYPPLGRFA VRDMRQTVAVGVIKAVDKKAAGAGKVTKSAQKAQKAK

>EF1G(bait) · .

aagtlytypenwrafkaliaaqysgaqvrvlsapphfhfgqtnrtpeflrkfpagkvpafeGddgf CVFESNAIAYYVSNEELRGSTPEAAAQVVQWVSFADSDIVPPASTWVFPTLGIMHHNKQATENAKE EVRRILGLLDAYLKTRTFLVGERVTLADITVVCTLLWLYKQVLEPSFRQAFPNTNRWFLTCINQPQ FRAVLGEVKLCEKMAQFDAKKFAETQPKKDTPRKEKGSREEKQKPQAERKEEKKAAAPAPEEEMDE CEOALAAEPKAKDPFAHLPKSTFVLDEFKRKYSNEDTLSVALPYFWEHFDKDGWSLWYSEYRFPEE LTQTFMSCNLIT@MFQRLDKLRKNAFASVILFGTNNSSSISGVWVFRGQELAFPLSPDWQVDYESY TWRKLDPGSEETQTLVREYFSWEGAFQHVGKAFNQGKIFK

>EF1G(prey) AAGTLYTYPENWRAFKALIAAQYSGAQVRVLSAPPHFHFGQTNRTPEFLRKFPAGKVPAFEGDDGF CVFESNAIAYYVSNEELRGSTPEAAAQVVQWVSFADSDIVPPASTWVFPTLGIMHHNKQATENAKE EVRRILGLLDAYLKTRTFLVGERVTLADITVVCTLLWLYKQVLEPSFRQAFPNTNRWFLTCINQPQ FRAVLGEVKLCEKMAQFDAKKFAETQPKKDTPRKEKGSREEKQKPQAERKEEKKAAAPAPEEEMDE CEQALAAEPKAKDPFAHLPKSTFVLDEFKRKYSNEDTLSVALPYFWEHFDKDGWSLWYSEYRFPEE LTQTFMSCNLITGMFQRLDKLRKNAFASVILFGTNNSSSISGVWVFRGQELAFPLSPDWQVDYESY TWRKLDPGSEETQTLVREYFSWEGAFQHVGKAFNQGKIFK

>FEZ1

GNCSDTEIHEKEEBEFNEKSENDSGINEEPLLTADQVIBEIBEMMQNSPDPBEBEEVLEBEDGGET SSOADSVLLQEMQALTQTFNNNWSYEGLRHMSGSELTELLDQVEGAIRDFSEELVQQLARRDELEF EKEVKNSFITVLIEVQNKQKEQRELMKKRRKEKGLSLQSSRIEKGNQMPLKRFSMEGISNILQSGI RQTFGSSGTDKQYLNTVIPYEKKASPPSVEDLQMLTNILFAMKEDNEKVPTLLTDYILKVLCPT

>G45IP1 ·

MASSGELGSLFDHHVQRAVCDTRAKYREGRRPRAVKVYTINLESQYLLIQGVPAVGVMKELVERF ALYGAIEQYNALDEYPAEDFTEVYLIKFMNLQSARTAKRKMDEQSFFGGLLHVCYAPEFETVEETR KKLQMRKAYVVKTTENKDHYVTKKKLVTEHKDTEDFRQDFHSEMSGFCKAALNTSAGNSNPYLPYS CELPLCYFSSKCMCSSGGPVDRAPDSSKDGRNHHKTMGHYNHNDSLRKTQINSLKNSVACPGAQKA ITSSEAVDRFMPRTTQLQERKRRREDDRKLGTFLQTNPTGNEIMIGPLLPDLSKVDMHDDSLNTTA NLTRHKEKEVTSSVPKPPEDKPEDVHTSHPLKQRRRI

>G45IP2

RTCMPYIFSLSLEALKCFRIRNNEKMLSDSHGVETIRDILPDTSLGGPSFFKIITAKAVLKLQAGN AEEAALWRDLVRKVLASYLETAEEAVTLGGSLDENCQEVLKFATRENGFLLQYLVAIPMEKGLDSQ GCFCAGCSRQIGFSFVRPKLCAFSGLYYCDICHQDDASVIPARIIHNWDLTKRPICRQALKFLTQI RAQPLINLQMVNASLYEHVERMHLIGRRREQLKLLGDYLGLCRSGALKELSKRLNHRNYLLESPHR FSVADLQQIADGVYEGFLKALIEFASQHVYHCDLCTQRGFICQICQHHDLIFPFEFDTTVRCAECK TVFHOSCQAVVKKGCPRCARRKYOEONIFA

>G45IP3

PNRGPLSPPNDLRPSHVISLPLHNAPHTRPTNQHTNHIPMMARCNTRKHIPRPPHTTCPKRPSIRD NPIYYLRSFFLRRIFLSLLPLQPSPYPPIRRALAPNRHHPAKSPRSPTPKHIRITRIRSINHLSSP SGADD45G

GAGAEPGLECGWSWGAKGVCRWPGLGSPPRPPAGSRSLRWLLRRMQGAGKALHELLLSAQRQGCLT AGVYESAKVLNVDPDNVTFCVLAAGEEDEGDIALQIHFTLIQAFCCENDIDIVRVGDVQRLAAIVG AGEEAGAPGDLHCILISNPNEDAWKDPALEKLSLFCEESRSVNDWVPSITLPE >GIT1

POMADRSROKCMSQSLDLSELAKAAKKKLQALSNRLFEELAMDVYDEVDRRENDAVWLATQNHSTL
VTERSAVPFLPVNPEYSATRNQGRQKLARFNAREFATLIIDILSEAKRRQQGKSLSSPTDNLELSL
RSQSDLDDQHDYDSVASDEDTDQEPLRSTGATRSNRARSMDSSDLSDGAVTLQEYLELKKALATSE
AKVQQLMKVNSSLSDELRRLQREIHKLQAENLQLRQPPGPVPTPPLPSERAEHTPMAPGGSTHRRD
RQAFSMYEPGSALKPFGGPPGDELTTRLQPFHSTELEDDAIYSVHVPAGLYRIRKGVSASAVPFTP
SSPLLSCSQEGSRHTSKLSRHGSGADSDYENTQSGDPLLGLEGKRFLELGKEEDFHPELESLDGDL
DPGLPSTEDVILKTEQVTKNIQELLRAAQEFKHDSFVPCSEKIHLAVTEMASLFPKRPALEPVRSS
LRLLNASAYRLQSECRKTVPPEPGAPVDFQLLTQQVIQCAYDIAKAAKQLVTITTREKKQ
>hADA3

KDVDALLKKSEAQHEQPEDGCPFGALTQRLLQALVEENIISPMEDSPIPDMSGKESGADGASTSPR NQNKPFSVPHTKSLESRIKEELIAQGLLESEDRPAEDSEDEVLAELRKRQAELKALSAHNRTKKHD LLRLAKEEVSRQELRQRVRMADNEVMDAFRKIMAARQKKRTPTKKEKDQAWKTLKERESILKLLDG

DAERQEALGIVRRIGTDTEAATEPAGATVPAAAAARIGTVGPQPPAMPRRKRNAGSSSDGTEDSD FSTDLEHTDSSESDGTSRSARVTRSSARLSQSSQDSSPVRNLQSFGTEEPAYSTRRVTRSQQQPT PVTPKKYPLRQTRSSGSETEQVVDFSDRETKNTADHDESPPRTPTGNAPSSESDIDISSPNVSHDE SIAKDMSLKDSGSDLSHRPKRRRFHESYNFNMKCPTPGCNSLGHLTGKHERHFSISGCFLYHNLSA DECKVRAQSRDKQIEERMLSHRQDDNNRHATRHQAPTERQLRYKEKVAELRKKRNSGLSKEQKEKY MEHRQTYGNTREPLLENLTSEYDLDLFRRAQARASEDLEKLRLQGQITEGSNMIKTIAFGRYELDT WYHSPYPLEYARLGRLYMCEFCLKYMKSQTILRRHMAKCVWKHPPGDEIYRKGSISVFEVDGKKNKIYCQNLCLLAKLFLDHKTLYYDVEPFLFYVMTEADNTGCHLIGYFSKEKNSFLNYNVSCILTMPQYMRQGYGKMLIDFSYLLSKVEEKVGSPERPLSDLGLISYRSYWKEVLLRYLHNFQGKEISIKEISQETAVNPVDIVSTLQALQMLKYWKGKHLVLKRQDLIDEWIAKEAKRSNSNKTMDPSCLKWTPPKGT

>HD1.7

>HDd1.0

>HDd1.3

PRLQLELYKEIKKNGAPRSLRAALWRFAELAHLVRPQKCRPYLVNLLPCLTRTSKRPEESVQETLA
AAVPKIMASFGNFANDNEIKVLLKAFIANLKSSSPTIRRTAAGSAVSICQHSRRTQYFYSWLLNVL
LGLLVPVEDEHSTLLILGVLLTLRYLVPLLQQQVKDTSLKGSFGVTRKEMEVSPSAEQLVQVYELT
LHHTQHQDHNVVTGALELLQQLFRTPPPELLQTLTAVGGIGQLTAAKEESGGRSRSGSIVELIAGG
GSSCSPVLSRKQKGKVLLGEEEALEDDSESRSDVSSSALTASVKDEISGELAASSGVSTPGSAGHD
IITEOPRSOHTAGGLSGSGQL

>HDexQ20

>HDexO51

ADTLQGHRDRFMEQFTKLKDLFYRSSNLQYFKRLIQIPQLPENPPNFLRASALSEHISPVVVIPAE
ASSPDSEPVLEKDDLMDMDASQQNLFDNKFDDIFGSSF3SDPFNFNSQNGVNKDEKDHLIERLYRE
ISGLKAQLENMKTESQRVVLQLKGHVSELEADLAEQQHLRQQAADDCEFLRAELDELRRQREDTEK
AQRSLSEIERKAQANEQRYSKLKEKYSELVQNHADLLRKNAEVTKQVSMARQAQVDLEREKKELED
SLERISDQGQRKTQEQLEVLESLKQELATSQRELQVLQGSLETSAQSEANWAAEFAELEKERDSLV
SGAAHREEELSALRKELQDTQLKLASTEESMCQLAKDQRKMLLVGSRKAAEQVIQDASTRP
>HIP11

VDLVTACDIRYCAQDAFFQVKEVDVGLAADVGTLQRLPKVIGNQSLVNELAFTARKMMADEALGSG LVSRVFPDKEVMLDAALALAAEISSKSPVAVQSTKVNLLYSRDHSVAESLNYVASWNMSMLQTQDL VKSVQATTENKELKTVTFSKL

>HIP13

PCCSEDTIPSQVSDYDYFSVSGDQEADQQEFDKSSTIPRNSDISQSYRRMFQAKRPASTAGLPTTL GPAMVTPGVATIRRTPSTKPSVRRGTIGAGPIPIKTPVIPVKTPTVPDLPGVLPAPPDGPEERGEH SPESPSVGEGPQGVTSMPSSMWSGQASVNPPLPGPKPSIPEEHRQAIPESEAEDQEREPPSATVSP GQIPESDPADLSPRDTPQGEDMLNAIRRGVKLKKTTTNDRSAPRFS

SHIPIS

IHMAPPYPNLNMIETFICQVCEETLAHSVDSLEQLTGIRMLRHLTMTIDYHTLIANYMSGFLSLLT TANARTKFHVLKMLLNLSENPAVAKKLFSAKALSIFVGLFNIEETNDNIQIVIKMFQNISNIIKSG KMSLIDDDFSLEPLISAFREFEELAKQLQAQIDNQNDPEVGQQS

>HIP16

DEEERNHRQMIKEAFAGDDVIRDFLKEKREAVEASKPKDVDLTLPGWGEWGGVGLKPSAKKRRRFL IKAPEGPPRKDKNLPNVIINEKRNIHAAAHQVRVLPYPFTHHWQFERTIQTPIGSTWNTQRAFQKL TTPKVVTKPGHIINPIKAEDVGYRSSSRSDLSVIQRNPKRITTRHKKQLKKCSVD >HIP2

MANIAVQRIKREFKEVLKSEETSKNQIKVDLVDENFTELRGB<u>IAGPPDTPYEGGRYQLEIKIPETY</u>
PENPPKYRFITKIWHPNISSVICATCLDILKDQWAAAMTLRTVLLSLQALLAAAEPDDPQDAVVAN
QYKQNPEMFKQTARLWAHVYAGAPVSSPEYTKKIENLCAMGFDRNAVIVALSSKSWDVETATELLL
SNX

>HIP5(bait)

FLKSILKKESKYEHGYLKALIINQSFKFGNQKAAAIRDSIELTKEKGAEIPKTIKKLRNFDETSNI ENNAENSHSLKNKTGTTQQHSQQPHIQSGAGSNIISVSTCAVNSADTKKSREDSISENVTTLGGSG ADHMPLNCFIPSGYNFAKHAWPASKKEESKIPVHDDSKTKQGKPQRGRAKIIRKPGSAKVQSGFIC TNRKGAVTQPQSASKVNIFTQAQGKLIIPCPPPQSTSNIRSGKNIQVSQCQPVTPENPQNIITHNS FNSKHVLPTEHSLNQWNQESSSPLSNACSDLVTVIPSLPSYCSSECQTFAKINHSNGTQAVARQDA TLYCTQRSPVCEESYPSVTLRTAEBESVPLWKRGPNVLHQNKRATGSTVMRRKRIAETKRRNILEQ KRQNPGSVGQKYSEQINNFGQSVLLSSSEPKQTTRGTSYIEEVSDSTSEFLMAENLVKASVPEDEI LTVLNSKQIQKSNLPLNKTQQFNICTLSAEEQKILESINDLNERLHYIQESICKNPSIKNTLQIIP LLEKREDRTSSCRDKR

>HIP5(prey)

FLKSILKKESKYEHGYLKALIINQSFKFGNQKAAAIRDSIELTKEKGAEIPKTIKKLRWFDETSNI ENNAENSHSLKNKTGTTQQHSQQFHIQSGAGSNIISVSTCAVNSADTKKSREDSISENVTTLGGSG ADHMPLNCFIPSGYNFAKHAWPASKKEESKIPVHDDSKTKQGKPQRGRAKIIRKPGSAKVQSGFIC TNRKGAVIQPQSASKVNIFTQAQGKLIIPCPPPPQSTSNIRSGKNIQVSQCQPVTPENPQNIITHNS FNSKHVLPTEHSLNQWNQESSSPLSNACSDLVTVIPSLPSYCSSECQTFAKINHSNGTQAVARQDA TLYCTQRSPVCEESYPSVTLRTAEEESVPLWKRGPNVLHQNKRATGSTVMRRKRIAETKRRNILEQ KRQNPGSVGQKYSEQINNFGQSVLLSSSEPKQTTRGTSYIEEVSDSTSEFLMAENLVKASVPEDEI LTVLNSKQIQKSNLPLNKTQQFNICTLSAEEQKILESLNDLNERLHYIQESICKNPSIKNTLQIIP LLEKREDRTSSCRDKR

>HMP

QEQVKIESLAKSLEDALRQTASVTLQAIAAQNAAVQAVNAHSNILKAAMDNSEIAGEKKSAQWRTV EGALKERRKAVDEAADALLKAKEELEKMKSVIENAKKKEVAGAKPHITAAEGKLHNMIVDLDNVVK KVQAAQSEAKVVSQYHELVVQARDDFKRELDSITPEVLPGWKGMSVSDLADKLSTDDLNSLIAHAH RRIDQLNRELAEQKATEKQHITLALEKQKLEEKRAFDSAVAKALEHHRSEIQAEQDRKIEEVRDAM ENEMRTQLRRQAAAHTDHLRDVLRVQEQELKSEFEQNLSEKLSEQELQFRRLSQEQVDNFTLDINT AYARLRGIEQAVQSHAVAEEEARKAHQLWLSVEALKYSMKTSSABTPTIPLGSAVEAIKANCSDNE FTQALTAAIPPESLTRGVYSEETLRARFYAVQKLARRVAMIDETRNSLYQYFLSYLQSLLLFPPQQ LKPPPELCPEDINTFKLLSYASYCIEHGDLELAAKFVNQLKGESRRVAQDWLKEARMTLETKQIVE ILTAYASAVGIGTTQVQPE

~HP28

PPADSLLKYDTPVLVSRNTEKRSPKARLLKVSPQQPGPSGSAPQPPKTKLPSTPCVPDPTKQAEEI LNAILPPREWVEDTQLWIQQVSSTPSTRMDVVHLQEQLDLKLQQRQARETGICPVRRELYSQCFDE LIREVTINCAERGLLLLRVRDEIRMTIAAYQTLYESSVAFGMRKALQAEQGKSDMERKIAELETEK RDLERQVNEQKAKCEATEKRESERRQVEEKKHNEEIQFLKRTNQQLKAQLEGIIAPKK >HSPC232

RRRADGCIYGVSRRARVVAYRDEMWSEGRYEYERIPRERAPPRSHPSDESGYRWTRDDHSASRQP EYRDMRDGFRRKSFYSSHYARERSPYKRDNTFFRESPVGRKDSPHSRSGSSVSSRSYSPERSKSYS FHQSQHRNKERPVQSLKTSRDTSPSSGSAVSSSKVLDKPSRLTEKELAEAASKWAAEKLEKSDESN LPEISEYEAGSTAPLFTDQPEEPESNTTHGIELFEDSQLTTRSKAIASKTKEIEQVYRQDCETFGM VVKMLIEKDPSLEKSIQFALRQNLHEIGERCVEELKHFIAEYDTSTQDFGEPF

>HYPA

GRRSSLSPTMRPGTGAERGGLMMGHPGMHYAPMGMHPMGQRANMPPVPHGMMPQMMPPMGGPPMG
QMPGMMSSVMPGMMSHMSQASMQPALPPGVNSMDVAAGTABGAKSMWTEHKSPDGRTYYYNTETK
QSTWEKPDDLKTPAEQLLSKCPWKEYKSDSGKPYYYNSQTKESRWAKPKELEDLEGYQNTIVAGSL
ITKSNLHAMIKAEESSKQEECTTTSTAPVPTTEIPTTMSTMAAAEAAAAVVAAAAAAAAAAAAAAAA
NASTSASNTVSGTVPVVPEPEVTSIVATVVDNENTVTLSTREQAQLTSTPALQDGSVEVSGNTGEE
TSKQETVADFTPKKEEEESQPAKKTYTWNTKEEAKQAFKELLKEKRVPSNASWEQAMKMIINDPRY
SALAKLSEKKQAFNAYKVQAKKKEKKKKKK

WZFF

HARFAEAECLAESHQHLSKESLAGNKPANAVLHKVLNQLEELLSDMKADVTRLPATLSRIPPIAAR LQMSERSILSRLASKGTEPHPTPAYPPGPYATPPGYGAAFSAAPVGALAAAGANYSQMPAGSFITA ATNGPPVLVKKEKEMVGALVSDGLDRKEPRAGEVICIDD

TWAL

 $\tt LKEGSPLEDLALLEALSEVVQNTENLKDEVYHILKVLFLFEFDEQGRELQKAFEDTLQLMERSLPEINTLTYQQNSATPVLGPNSTANSIMASYQQQKTSVPVLDAELFIPPKINRRTQWKLSLLD$

- TMDD9

DFLILPGYIDFTADQVDLTSALTKKITLKTPLVSSPMDTVTEAGMAIAMALTGGIGFIHHNCTPEF
QANEVRKVKKYEQGFITDPVVLSPKDRVRDVFEAKARHGFCGIPITDTGRMGSRLVGIISSRDIDF
LKEEEHDCFLEEIMTKREDLVVAPAGITLKEANEILQRSKKGKLPIVNEDDELVAIIARTDLKKNR
DYPLASKDAKKQLLCGAAIGTHEDDKYRLDLLAQAGVDVVVLDSSQGNSIFQINMIKYIKDKYPNL
QVIGGNVVTAAQAKNLIDAGVDALRVGMGSGSICITQEVLACGRPQATAVYKVSEYARRFGVPVIA
DGGIQNVGHIAKALALGASTVMMGSLLAATTEAPGEYFFSDGIRLKKYRGMGSLDAMDKHLSSQNR
YFSEADKIKVAQGVSGAVQDKGSIHKFVPYLIAGIQHSCQDIGAKSLTQVRAMMYSGELKFEKRTS
SAQVEGGVHSLHSYEKRLF

>KPNA2

AWALTNIASGTSEQTKAVVDGGAIPAFISLLASPHAHISEQAVWALGNIAGDGSVFRDLVIKYGAV
DPLLALLAVPDMSSLACGYLRNLTWTLSNLCRNKNPAPPIDAVEQILPTLVRLLHHDDPEVLADTC
WAISYLTDGPNERIGMVVKTGVVPQLVKLLGASELPIVTPALRAIGNIVTGTDEQTQVVIDAGALA
VFPSLLTNPKTNIQKEATWTMSNITAGRQDQIQQVVNHGLVPFLVSVLSKADFKTQKEAVWAVTNY
TSGGTVEQIVYLVHCGIIEPLMNLLTAKDTKIILVILDAISNIFQAAEKLGETEKLSIMIEECGGL
DKIEALQNHENESVYKASLSLIEKYFSVEEEEDQNVVPETTSEGYTFQVQDGAPGTFNF
>KPNB1

LAAVGLVGDLCRALQSNIIPFCDEVMQLLLENLGNENVHRSVKPQILSVFGDIALAIGGEFKKYLE VVLNTLQQASQAQVDKSDYDMVDYLNELRESCLEAYTGIVQGLKGDQENVHPDVMLVQPRVEFILS FIDHIAGDEDHTDGVVACAAGLIGDLCTAFGKDVLKLVEARPMIHELLTEGRRSKTNKAKTLATWA TKELRKLKNQA

>Ku70

KTRTFNTSTGGLLLPSDTKRSQIYGSRQIILEKEETEELKRFDDPGLMLMGFKPLVLLKKHHYLRP SLFVYPEESLVIGSSTLFSALLIKCLEKEVAALCRYTPRRNIPPYFVALVPQEEELDDQKIQVTPP GFQLVFLPFADDKRKMPFTEKIMATPEQVGKMKAIVEKLRFTYRSDSFENPVLQQHFRNLEALALD LMEPEQAVDLTLPKVEAMNKRLGSLVDEFKELVYPPDYNPEGKVTKRKHDNEGSGSKRPKVEYSEE ELKTHISKGTLGKFTVPMLKEACRAYGLKSGLKKQELLEALTKHFQD

STJIC7B1

VDAVAVDAAAVSAKAEKVHELNEKIGKLLAKAEQLGAEGNVDESQKILMEVEKVRAKKKEAEEEYR NSMPASSFQQQKLRVCEVCSAYLGLHDNDRRLADHFGGKLHLGFIQIREKLDQLRKTVAEKQEKRN QDRLRREEREREERLSRRSGSRTRDRRRSRSRDRRRRSSRSTSRERRKLSRSRSRDRHRRHRSRS RSHSRGHRRASRDRSAKYKFSRERASREESWESGRSERGPPDWRLESSNGKMASRRSEEKEAGEI

>MAGEH1

ASFPRTAVSFEPLAGDMPRGRKSRRRRNARAAEENRNNRKIQASEASETPMAASVVASTPEDDLSG PEEDPSTPEEASTTPEEASSTAQAQKPSVPRSNFQGTKKSLLMSILALIFIMGNSAKEALVWKVLG KLGMQPGRQHSIFGDPKKIVTEEFVRRGYLIYKPVPRSSPVEYEFFWGPRAHVESSKLKVMHFVAR VRNRCSKDWPCNYDWDSDDDAEVEAILNSGARGYSAP

>MAP11c3

QRSFMDRKKEYQOTRDQHPSKIPYTIERYKGEKQLPVLDKTKPLVPDHVNMSELVKIIRRRLQLN PTQAFFLLVNQHSMVSVSTPIADIYEQEKDEDGFLYMVYASQETFGF

PKEQVQSGAGDGTGSGDPAAGTFTTQPAVGPAPEPSAEPKPAPAQGTGSGQKSGSRTKTGSFCRSM IIGDSDAPWTRYVFQGPYGPRATGLGTGKAEGIWKTPAAYIGRRPGVSGPERAAFIRELQEALCPN PPPTKKITEDDVKVMLYLLEEKERDLNTAARIGQSLVKQNSVLMEENNKLETMLGSAREEILHLRK QVNLRDDLLQLYSDSDDDDDEEDEEDEEGEEEEREGQRDQDQQHDHPYGAPKPHPKAETAHRCPQ LETLQQKLRLLEEENDHLREEASHLDNLEDEEQMLILECVEQFSEASQQMAELSEVLVLRLEGYER QQKEITQLQAEITKLQQRCQSYGAQTEKLQQMLASEKGIHSESLRAGSYMQDYGSRPRDRQEDGKS HRQRSSMPAGSVTHYGYSVPLDALPSFPETLAEELRTSLRKFITDPAYFMERRDTHCREGRKKEQR AMPPPPAX

>mp53 °

VTETPGPVAPAPATPWPLSSFVPSQKTYQGNYGFHLGFLQSGTAKSVMCTYSPPLNKLFCQLAKTC PVQLWVSATPPAGSRVRAMAIYKKSQHMTEVVRRCPHHERCSDGDGLAPPQHLIRVEGNLYPEYLE DRQTFRHSVVVPYEPPEAGSEYTTIHYKYMCNSSCMGGMNRRPILTTITLEDSSGNLLGRDSFEVR VCACPGRDRRTEEENFRKKEVLCPELPPGSAKRALPTCTSASPPQKKKPLDGEYFTLKIRGRKRFE MFRELNEALELKDAHATEESGDSRAHSSYLKTKKGQSTSRHKKTMVKKVGPDSD

RDRVENEAEKDLQCHAPVRLDLPFEKPLTSSLAKQEEVEQTPLQEALNQLMRQLQRKDPSAFFSFP VTDF1APGYSMIIKHPMDFSTMKEKIKNNDYQSIEELKDNFKLMCTNAMIYNKPETIYYKAAKKLL HSGMKILSQERIQSLKQSIDFMADLQKTRKQKDGTDTSQSGEDGGCWQREREDSGDAEAHAFKSPS KENKKKDKDMLEDKFKSNNLEREQEQLDRIVKESGGKLTRRLVNSQCEFERRKPDGTTTLGLLHPV DPIVGEPGYCPVRLGMTTGRLQSGVNTLQGFKEDKRNKVTPVLYLNYGPYSSYAPHYDSTFANISK DDSDLIYSTYGEDSDLPSDFSIHEFLATCQDYPYVMADSLLDVLTKGGHSRTLQEMEMSLPEDEGH TRILDTAKEMEITEVEPPGRLDSSTQDRLIALKAVTNFGVPVEVFDSEEAEIFQKKLDETTRLLRE LQEAQNERLSTRPPPNMICLLGPSYREMHLAEQVTNNLKELAQQVTPGDIVSTYGVRKAMGISIPS PVMENNFVDLTEDTEEPKKTDVAECGPGGS

>NEFL

LSPLSSLSGLPPPPRAGEPPAATMSSFSYEPYYSTSYKRRYVETPRVHISSVRSGYSTARSAYSSY SAPVSSSLSVRRSYSSSSGSLMPSLENLDLSQVAAISNDLKSIRTQEKAQLQDLNDRFASFIERVH ELEQQNKVLEAELLVLRQKHSEPSRFRALYEQEIRDLRLAAEDATNEKQALQGEREGLEETLRNLQ ARYEEEVLSREDAEGRLMEARKGADEAALARAELEKRIDSLMDEISFLKKVHEEEIAELQAQIQYA QISVEMDVTKPDLSAALKDIRAQYEKLAAKNMQNAEEWFKSRFTVLTESAAKNTDAVRAAKDEVSE SRRLLKAKTLEIEACRGMNEALEKQLQELEDKQNADISAMQDTINKLENELRTTKSEMARYLKEYQ DLLNVKMALDIEIAAYRKLLEGEETRLSFTSVGSITSGYSQSSQVFGRSAYGGLQTSSYLMSTRSF PSYYTSHVQEEQIEVEETIEAAKAEEAKDEPPSEGEAEEEEKDKEEAEEEEAAEEEEAAKEESEEA KEEEEGGEGEEGEETKEAEEEEKKVEGAGEEQAAKKKD

>p53

MEEPQSDPSVEPPLSQETFSDLWKLLPENNVLSPLPSQAMDDLMLSPDDIEQWFTEDPGPDEAPRM PEAAPPVAPAPAPAPAPAPAPAPSWPLSSSVPSQKTYQGSYGFRLGFLHSGTAKSVTCTYSPALNK MFCQLAKTCPVQLWVDSTPPPGTRVRAMAIYKQSQHMTEVVRRCPHHERCSDSDGLAPPQHLIRVE GNLRVEYLDDRNTFRHSVVVPYEPPEVGSDCTTIHYNYMCNSSCMGGMNRRPILTIITLEDSGNLLGRNSFEVRVCACPGRDRRTEEENLRKKGEPHHELPPGSTKRALPNNTSSSPQPKKKPLDGEYFTLQIRGRERFEMFRELNEALELKDAQAGKEPGGSRAHSSHLKSKKGQSTSRHKKLMFKTEGPDSD

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>PFN2

APRRPRCSAKGSKMAGWQSYVDNLMCDGCCQEAAIVGYCDAKYVWAATAGGVFQSITPIEIDMIVG KDREGFFTNGLTLGAKKCSVIRDSLYVDGDCTMDIRTKSQGGEPTYNVAVGRAGRVLVFVMCKEGV HGGGLNKKAYSMAKYLRDSGF

>PIASy(bait)

LVEAKNMYMSFRYSDLOMLLGFYGRSKSGLKHELYFRALGENOFDCSPELFKKEKELYBTRYAGKTV SEPAPOPHRPDDPLTMHSTYDRAGAVPRTPLAGPNIDYPVLYGKYLNGLGRLPAKTLKPEVRLVKL PFFNMLDELLKPTELVPQNNEKLQESPCIFALTPRQVELIRNSRELQPGVKAVQVVLRICYSDTSC PQEDQYPPNIAVKVNHSYCSVPGYYPSNKPGVEPKRPCRPINLTHLMYLSSATNRITVTWGNYGKS YSVALYLVRQLTSSELLQRLKTIGVKHPELCKALVKEKLRLDPDSEIATTGVRVSLICPLVKMRLS VPCRAETCAHLQCFDAVFYLQMNEKKPTWMCPVCDKPAPYDQLIIDGLLSKILSECEDADEIEYLV DGSWCPIRAEKERSCSPQGAILVLGPSDANGLLPAPSVNGSGALGSTGGGGPVGSMENGKPGADVV DLTLDSSSSSEDEEEEEEEEEDEDEEGPRPKRRCPFQKGLVPAC >PIASy(prey)

LVEAKNMVMSFRVSDLQMLLGFVGRSKSGLKHELVTRALQLVQFDCSPELFKKIKELYETRYAKKN SEPAPQPHRPLDPLTMHSTYDRAGAVPRTPLAGPNIDYPVLYGKYLNGLGRLPAKTLKPEVRLVKL PFFNMLDELLKPTELVPQNNEKLQESPCIFALTPRQVELIRNSRELQPGVKAVQVVLRICYSDTSC PQEDQYPPMIAVKVNHSYC5VPGYYP5NKPGVEPKRPCRPINLTHLMYLSSATNRITVTWGNYGKS YSVALYLVRQLTSSELLQRLKTIGVKHPELCKALVKEKLRLDPDSEIATTGVRVSLICPLVKMRLS VPCRAETCAHLQCFDAVFYLQMNEKKPTWMCPVCDKPAPYDQLIIDGLLSKILSECEDADEIEYLV DGSWCPTRAEKERSCSPQGATLVLGPSDANGLLPAPSVNGSGALGSTGGGGPVGSMENGKPGADVV DLTLDSSSSSEDEEEEEEEEEDEDEEGPRPKRRCPFQKGLVPAC . >PLIP

GEIIEGCRLPVLRRNQDNEDEWPLAEILSVKDISGRKLFYVHYIDFNKRLDEWVTHERLDLKKIQF PKKEAKTPTKNGLPGSRPGSPEREVKRKVEVVSPATFVPSETAPASVFPQNGAARRAVAAQPGRKR KSNCLGTDEDSQDSSDGIPSAPRMTGSLVSDRSHDDIVTRMKNIECIELGRHRLKPWYFSPYPQEL TTLPVLYLCEFCLKYGRSLKCLQRHLTKCDLRHPPGNEIYRKGTISFFEIDGRKNKSYSQNLCLLA KCFLDHKTLYYDTDPFLFYVMTEYDCKGFHIVGYFSKEKESTEDYNVACILTLPPYQRRGYGKLLI EFSYELSKVEGKTGTPEKPLSDLGLLSYRSYWSQTILEILMGLKSESGERPQITINEISEITSIKK EDVISTLQYLNLINYYKGQYILTLSEDIVDGHERAMLKRLLRIDSKCLHFTPKDWSKRGKW >PTN

L5QRQDQVPRLPVQKSRQESPRAEENPKWREGKKETSESSVQKAGRAAAAQAGAAASRVPGLSGSN LAPCNKGRLSAREDVSNSKMQAQQYQQQRRKFAAAFLAFIFILAAVDTAEAGKKEKPEKKVKKSDC GEWQWSVCVPTSGDCGLGTREGTRTGAECKQTMKTQRCKIPCNWKKQFGAECKYQFQAWGECDLNT ALKTRTGSLKRALHNAECQKTVTISKPCGKLTKPKPQAESKKKKKEGKKQEKMLD

SNYINAALMDSYRQPAAFIVTQYPLPNTVKDFWRLVYDYGCTSIVMLNEVDLSQGCPQYWPEEGML RYGPIQVECM3CSMDCDVINRIFRICNLTRPQEGYLMVQQFQYLGWASHREVPGSKRSFLKLILQV EKWQEECEEGEGRTIIHCLNGGGRSGMFCAIGIVVEMVKRQNVVDVFHAVKTLRNSKPNMVEAPEQ · YRFCYDVALEYLESS

>SETBD1

KASTSGLG_KDEGDIKQAKKEDTDDRNKMSVVTESSRNYGYNPSPVKPEGLRRPPSKTSMHQSRRL MASAQSNPDDVLTLSSSTESEGESGTSRKPTAGQTSATAVDSDDIQTISSGSEGDDFEDKKNMTGP MKRQVAVKSTRGFALKSTHGIAIKSTNMASVDKGESAFVRKNTRQFYDGEESCYIIDAKLEGNLGR YLNHSCSPNLFVQNVFVDTHDLRFPWVAFFASKRIRAGTELTWDYNYEVGSVEGKELLCCCGAIEC RGRLL .

>SH3GL3

VAGLKKOFHKASOLFSEKISGAEGTKLDDEFLDMERKIDVTNKVVAEILSKTTEYLOPNPAYRAKL GMLNTVSKIRGOVKTTGYPOTEGLIGDCMLKYGKELGEDSTFGNALIEVGESMKLMAEVKDSLDIN VKÖTFIDPLOLLODKDLKEIGHHLKKLEGRRLDYDYKKKRVGKIPDEEVROAVEKFEESKELAERS MFNFLENDVEOVSQLAVFIEAALDYHROSTEILOELOSKLOMRISAASSVPRREYKPRFVKRSSSE LNGVETTESVARTTESNIPMOORGESLYDFEPENOOFLGFKEEDELTETNOTDENWYEGVEHGESG

FFPINYVEVIVPLPQ

>SUMO-2

RPRAQÜRRESGGAESVTRPLRAASPAPPPRAARAAMSEEKPKEGVKTENDHINLKVÄGQDGSVVQF KIKRHTPLSKLMKAYCERQGLSMRQIRFRFDGQPINETDTPAQLEMEDEDTIDVFQQQTGGVPËSS LAGHSF

>SÜMO-3

PSSTAAASFFCRSWCCLCARLVRTWYLFCEAAAEETPALAMADEKPKEGVKTENNDHINLKVAGQD GSVVQFKIKRHTPLSKLMKAYCERQGLSMRQIRFRFDGQPINETDTPAQLEMEDEDTIDVFQQQTG GVY

>TAL1

SSPVKRQRMESALDOLKOFTTVVADTGDFHAIDEYKPQDATTNPSLILAAAQMPAYQELVEEAIAY GRKLGGSQEDQIKNAIDKLFVLFGAEILKKIPGRVSTEVDARLSFDKDAMVARARRLIELYKEAGI SKDRILIKLSSTWEGIQAGKELEEQHGIHCNMTLLFSFAQAVACAEAGVTLISPFVGRILDWHVAN TDKKSYEPLEDPGVKSVTKIYNYYKKFSYKTIVMGASFRNTGEIKALAGCDFLTISPKLLGELLQD NAKLVPVLSAKAAQASDLEKIHLDEKSFRWLHNEDQMAVEKLSDGIRKFAADAVKLERMLTERMFN AENGK

>TCPG

QTDIEITREEDFTRILOMEEEYIQQLCEDIIQLKPDVVITEKGISDLAQHYLMRANITAIRRVRKT DNNRIARACGARIVSRPEELREDDVGTGAGLLEIKKIGDEYFTFITDCKDPKACTILLRGASKEIL SEVERNLODAMQVCRNVLLDPQLVPGGGASEMAVAHALTEKSKAMTGVEQWPYRAVAQALEVIPRT LIQNCGASTIRLLTSLRAKHTQENCETWGVNGETGTLVDMKEIGÏWEPLAVKLQTYKTAVETÄVLL LRIDDIVSGHKKKGDDQSRQGGAPDAGQE

>VIM

SPRORRSRAPTTHTHRALVRLFSGSQSAPPPPPRPSPPSAMSTRSVSSSSYRRMFGGPGTASRPS SSRSYVTTSTRTYSLGSALRPSTSRSLYASSPGGVYATRSSAVRLRSSVPGVRLLQDSVDFSLADA INTEFKNTRTNEKVELQELNDRFANYIDKVRFLEQQNKTLLAELEQLKGQGKSRLGDLYEEEMREL RRQVDQLTNDKARVEVERDNLAEDIMRLREKLQEEMLQREEAENTLQSFRQDVDNASLARLDLERK VESLQEETAFLKKLHEEETQELQAQIQEQHVQIDVDVSKPDLTAALRDVRQQYESVAAKNLQEAEE WYKSKFADLSEAANRINDALRQAKQESTEYRRQVQSLTCEVDALKGTNESLERQMREMEENFAVEA ANYODTIGRLQDEIQNMKEEMARHLREYQDLLNVKMALDIEIATYRKLLEGEESRISLPLPNFSSL NLRETNLDSLPLVDTHSKRTLLIKTVETRDGQVINETSQHHDDLE

>VIMC

QEEMLOREEAENTLOSFRODVDNASLARLDLEKKVESLOEEIAFLKKLHEEBIQELOAQIQEQHVQ IDVDVSKPDLTAALRDVRÖQYESVÄÄKNLOEAEEWYKSKFADLSEAANRNNDALROAKQESTEYRR QVQSLTCEVDALKGTNESLERQMREMEENFÄVEAANYQDTIGRLQDEIQNMKEEMARHLREYQDLL NVKMALDIEIATYRKLLEGEESRISLFLPNFSSLNLRETNLDSLPLVDTHSKRTLLIKTVETRDGQ VINETSOHHDDLE

>ZHX1
EQTINDLTFDGSFVKEENAEQAESTEVSSSGISISKTPIMKMMKNKVENKRIAVHHNSVEDVPEEK
ENEIKPDREEIVENPSSSASESNTSTSIVNRIHPSTASTVVTPAAVLPGLAQVITAVSAQQNSNLI
PKVLIPVNSIPTYNAALDNNPLLLNTYNKFPYPTMSEITVLSAQAKYTEEQIKIWFSAQRLKHGVS
WTPEEVEEARRKQFNGTVHTVPQTITVIPTHISTGSNGLPSILQTCQIVGQPGLVLTQVAGTNTLP
VTAPIALTVAGVPSQNNIOKSQVPAAQPTAETKPATAAVPTSQSVKHETALVNPDSFGIRAKKTKE
QLAELKVSYLKNQFPHDSEIIRLMKITGLTKGEIKKWFSDTRYNQRNSKSNQCLHLNNDSSTTIII
DSSDETTESPTVGTAQPKQSWNPFPDFTPQKFKEKTAEQLRVLQASFLNSSVLTDEELNRLRAQTK
LTRREIDAWFTEKKKSKALKEEKMEIDESNAGSSKEEAGETSPADESGAPKSGSTGKICKKTPEQL
HMLKSAFVRTQWPSPEEYDKLAKESGLARTDIVSWFGDTRYAWKNGNLKWYYYYQSANSSSMNGLS
SLRKRGRGRPKGRGRGRPRGRPRGSKRINNWDRGPSLIKFKTGTAILKDYYLKHKFLNEQDLDELV
NKSHMGYEQVREWFAERQRRSELGIELFEENEEEDEVIDDQEEDEEETDDSDTWEPPRHVKRKLSK

>ZNF33B

CYECGKTFCLKSDLTIHQRTHTGEKPFACPECGKFFSHKSTLSQHYRTHTGEKPYECHECGKIFYN KSYLTKHNRTHTGEKPYECNECGKTFCQKSQLTQHQRIHIGEKPYECNECGKAFCHKSALIVHQRT HTQEKPYKCNECGKSFCVKSGLILHERKHTGEKPYECNECGKSFSHKSSLTVHYRAHTGEKSCQCN ECGKIFYRKSDLAKHQRSHTGEKPYECNTCRKTFSQKSNLIVHQRTHIGEKPYE

>ALEX2 GCCGAATCAGTAGTTGGGGCTGCAATGGCTTCTGCAATAGCACCACCTCCCGGGGTGACACACGCC CTTGGGGCTGCAGAGCCCCTGCAATGGCAGGGGCTCCCAAAGTGGCAGAAGCTCCCAGAGAAGCG GAGACTTCCAGGGCAGCGGTGCCTCCTGGGACAGTGGTGCCTACCGAAGCGGCAGCACCCACTGAG GTGACCGAGGGTCCTGGGGTAGCAGCACCTACCAAGGTAGCTGAAGCTCCCGGGGTGGCATCGCCT ACCGAGGCAGCTGAGGCTCCTGTGCCGGCAACGCCTACTGGGGCTGCAGCACC<u>TACTGGGGCTGCA</u> GAGTCTCCTGGAACTTCTGGTTCCCCTAGAACAGCGGTGGTTCCTGGAACATCAGCTGCCAAGAAA GCAACCCCTGGGGCTCACACTGGGGCTATACCGAAAGCCACATCAGCGACTGGAGCGGTACCCAAA GGTGGAGGCAAGGGTGTAACCAGGTCCCGGAATGGGGGCAAGGGGCAAGAGAAAAGCAAAAGTT GAAGTAGACGAACTGGGGATGGGCTTCCGTCCTGGAGATGGGGCTGCAGCAGCTGCTGCAGCCTCT GCTAATGGCGGACAGGCTTTCCTGGCAGAGGTCCCTGATTCTGAGGAAGGGGAGTCCGGGTGGACT GCCATGCAGAAGCGCCCCTTTCCTTATGAAAITGATGAGATTCTGGGTGTCCGCGATCTCAGGAAG GTCCTTGCCTTGCTTCAGAAATCTGATGATCCTTTCATCCAACAGGTAGCTTTGCTCACTCTGAGC AACAATGCCAATTATTCATGCAATCAAGAGACAATCCGCAAATTGGGAGGCCTCCCAATTATTGCA AACATGATCAACAAAACTGATCCACACATTAAGGAAAAAGCCTTAATGGCCATGAATAACCTGAGT GACTACCAACACCTGCTTGTCAATTCCATTGCAAACTTTTTCCGTTTGCTATCTCAGGGAGGTGGA AAAATCAAGGTTGAGATTTTGAAAATCCTTTCGAATTTTGCTGAAAATCCAGATATGTTGAAGAAA $\verb|CTTCTCAGTACCCAAGTGCCAGCATCATTTAGTTCCCTCTATAATTCTTACGTGGAATCAGAAATC| \\$ CTTATTAATGCCCTTACTCTATTTGAGATTATCTATGACAATCTCAGAGCAGAGTGTTTAACTAT AGAGCCTTAGCAAATCACCATGACCTCTTAGTGAAAGTGAAAGTTATAAAACTAGTGAACAAATTC

GAGGAAGAGGAATCCTTCCCACAGCCAGTAGATGATTACTTCGTGGAGCCTCCGCAGGCTGAA GAGGAAGAGGAAACGGTCCCACCCCCAAGCTCCCATACACTTGCAGTGGTCGGCAAAGTCACTCCC ACCCCGAGGCCCACAGACGGTGTGGATATTTACTTTGGCATGCCTGGGGAAATCAGTGAGCACGAG · GGGTTCCTGAGGGCCAAGATGGACCTGGAGGAGCGTAGGATGCGCCAGATTAATGAGGTGATGCGT CACTTCCAGTCCATTCTGCAGACTCTGGAGCGAGCGGTGTCTGGTGAGCGACAGCGCCTGGTGGAA ACCCACGCCACCGCGTCATCGCCCTTATCAACGACCAGCGCCGGGCTGCCTTGGAGGGCTTCCTG GCAGCCTGCAGGCAGATCCGCCTCAGGCGGAGCGTGTCCTGTTGGCCCTGCGGCGCTACCTGCGT GCGGAGCAGAAGGAACACACGCCACCACCACCACCATGTGGCCCCCTGGATCCCGAG AAGGCACAGCAGATGCGCTTCCAGGTGCATACCCACCTTCAAGTGATTGAGGAGAGGGTGAATCAG AGCCTGGGCCTGCTTGACCAGAACCCCCACCTGGCTCAGGAGCTGCGGCCCCAAATCCAGGAACTC CTCCACTCTGAACACCTGGGTCCCAGTGAATTGGAAGCCCCTGCCCCTGGGGGCAGCAGCAGCAGCAC AAGGGTGGGCTGCAGCCTCCAGATTCCAAGGATGACCCCCCATGACCCTTCCAAAAGGGTCCACA GAACAAGATGCTGCATCCCCTGAGAAAGAGAAGATGAACCCGCTGGAACAGTATGAGCGAAAGGTG ${\tt AATGCGTCTGTTCCAGGGGTTTCCCTTTCCACTCATCGGAGATTCAGAGGGATGAGCTGGCACCAG}$ CTGGGACAGGGTGTCCCGTGAGGCTGTGTCGGGTCTGC

SBAIPI
CGGCCGCGGACGAAGATGGCGACCGCCATGTACTTGGAGCACTATCTGGACAGTATCGAGAACCTT
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GCAGAGATTGACATCCTGGCTGCAGAGTACATCTCCACGGTGAAGACGCGTGTCTCCAGACCAGCGC
GTGGAGCGCCTGCAGAAGATCCAGAACGCCTACAGCAAGTGCAAGGAATACAGTGACGACAAAGTG
CAGCTGGCCATGCAGAACATCGTAGGAAACAACAACATTCGAAGGCTTGATGCAGACCTTGGCG
CGCTTTGAAGCAGATCTGAAGGACAAGATGGAGGCCAGCGATTTTGAAAGCTCCGGAGGGCGAGCG
TTAAAAAAAGGCCGGGGTCAGAAAGAAAAAAAGAGGGTCCGGGGGCCGAGGCAGGAGAACAACAAGAGAAAAAAAGAGGCTCCGGGGCCGAGGCAGCCATCCTGTCCGTG
CACCCCTCTGATGTGCTGGACATGCCCGTGGACCCAAACGAACCCACTTTGACTTTCACTTTGCC
TGCGTGGACCTTACCGCGAAACCAAAGGAAGAAAAATGG

>BAIP2

AGCCAGCAGGCCAGCGTGACCATGCACGATGTGGACGCCGAGTCCTTCGAGGTGTTGGTCGACTAC TGCTACACGGGTCGTGTGTCTCTCAGTGAGGCCAATGTGCAGCGCCTGTACGCGGCCTCCGACATG CTACAGCTGGAATATGTGCGGGAAGCCTGTGCCTCCTTCTTAGCCCGACGTCTTGACCTGACCAAC TGCACCGCCATCCTCAAGTTTGCAGACGCCTTCGACCATCACAAGCTTCGATCTCAGGCCCAGTCC TACATAGCTCACAACTTCAAGCAGCTCAGCCGAATGGGTTCAATTCGGGAGGAGACTCTAGCAGAT CTAACCCTGGCCCAGCTGCTGGCTGTCCTACGCCTGGATAGTCTGGACATAGAGAGTGAGCGGACT GTATGCCATGTAGCTGTGCAGTGGCTGGAGGCTGCCAAAGAGCGGGGTCCCAGTGCTGCAGAA GTCTTCAAGTGCGTGCGCTGGATGCACTTCACTGAAGAAGATCAGGACTACTTAGAAGGGCTGCTG ACCAAGCCCATCGTGAAGAAGTACTGCCTGGACGTTATTGAAGGGGCCCTGCAGATGCGCTATGGT GACCTGTTGTACAAGTCTCTGGTGCCAGTGCCAAACAGCAGCAGCAGCAGCAGCAGCAGCAACTCT CTTGTATCTGCAGCAGAAAATCCACCCCAGAGACTGGGTATGTGTGCCAAGGAGATGGTGATCTTC TTTGGACATCCTAGAGATCCCTTTCTCTGCTATGACCCTTACTCGGGGGACATTTACACAATGCCA CATGACATCTATCTAGCTGCTCAGCCCAGGAAAGACCTCTGGGTGTATAAACCAGCTCAGAATAGT TGGCAGCAACTTGCAGATCGCTTGCTGTGTCGTGAGGGCATGGATGTGGCATATCTCAATGGCTAC GTTCAGAGAAACCAGTGGGCATTGGTGGCTCCTGTCCCTCATTCCTTCTATTCCTTTGAACTCATA GTGGTTCAGAACTATCTTTATGCTGTCAACAGTAAGCGCATGCTTTGCTATGATCCTAGCCACAAT ATGTGGCTGAACTGTGCTTCTCTTAAACGTAGTGACTTTCAGGAAGCATGTGTCTTCAATGATGAA ATCTATTGTATCTGTGACATCCCAGTCATGAAGGTCTACAACCCAGCTAGGGGGAGAATGGAGGCGG ATTAGTAATATTCCTTTGGATTCAGAGACCCACAACTACCAGATTGTCAATCATGACCAAAAGTTG CTTCTCATCACTTCTACAACCCCACAATGGAAAAAGAACCGAGTGACAGTGTATGAGTATGATACT ${\tt AGGGAAGATCAGTGGATTAATATAGGTACCATGTTAGGCCTTTTGCAGTTTGACTCTGGCTTTATT}$ TGCCTTTGTGCTCGTGTTTATCCTTCCTGCCTTGAACCTGGTCAGAGTTTTATTACTGAGGAAGAT CATGCACGCAGTGAGTCTAGTACTGAATGGGACTTAGATGGATTCAGTGAGCTGGACTCTGAGTCA GGAAGTTCAAGTTCTTTTTCAGATGATGAAGTCTGGGTGCAAGTAGCACCTCAGCGAAATGCACAG GATCAGCAGGGTTCTTTG

Figure 6 (continued)

>BAIP3

TTGGCCGGTTTCGAGTCGCTGACCTGCAGCTTCCCTGTGGTTTCCCGAGGCCTCCTTGCTTCCCGC TCTCCGAGGAGCCTTTCATCCGAAGGCGGGACGATGCCGGATAATCGGCAGCCGAGGAACCGGCAG CCGAGGATCCGCTCCGGGAACGAGCCTCGTTCCGCGTCCGCCATGGAACCGGATGGTCGCGGTGCC TGGGCCCACAGTCGCGCGCGCTCGACCGCCTGGAGAAGCTGCTGCGCGTCGCGTTGTACTAAC ATTCTGAGAGAGCCTGTGTGTTTAGGAGGATGTGAGCACATCTTCTGTAGTAATTGTGTAAGTGAC TGCATTGGAACTGGATGTCCAGTGTTTACACCCCGGCCTGGATACAAGACTTGAAGATAAATAGA CAACTGGACAGCATGATTCAACTTTGTAGTAAGCTTCGAAATTTGCTACATGACAATGAGCCGTCA GATTTGAAAGAAGATAAACCTAGGAAAAGTTTGTTTAATGATGCAGGAAACAAGAAGAATTCAATT AAAATGTGGTTTAGCCCTCGAAGTAAQAAAGTCAGATATGTTGAGTAAAGCTTCAGTGCAAACC CAGCCTGCAATAAAAAAAGATGCAAGTGCTCAGCAAGACTCATATGAATTTGTTTCCCCAAGTCCT ACTTTAGCTGAAATCAACCAAAAATGGAATTTAGAGGCAGAAAAAGAAGATGGTGAATTTGACTCC AAAGAGGAATCTAAGCAAAAGCTGGTATCCTTCTGTAGCCAACCATCTGTTATCTCCAGTCCTCAG ATAAATGGTGAAATAGACTTACTAGCAAGTGGCTCCTTGACAGAATCTGAATGTTTTGGAAGTTTA ACTGAAGTCTCTTTACCATTGGCTGAGCAAATAGAGTCTCCAGACACTAAGAGCAGGAATGAAGTA GTGACTCCTGAGAAGGTCTGCAAAAATTATCTTACATCTAAGAAATCTTTGCCATTAGAAAATAAT GGAAAACGTGGCCATCACAATAGACTTTCCAGTCCCATTTCTAAGAGATGTAGAACCAGCATTCTG AGCACCAGTGGAGATTTTGTTAAGCAAACGGTGCCCTCAGAAAATATACCATTGCCTGAATGTTCT TCACCACCTTCATGCAAACGTAAAGTTGGTGGTACATCAGGGAGCAAAACAGTAACATGTCCGATG AATTCATTAGTCTTTCACCAGGTACACCACCTTCTACAT

CAACAATTCATTCCTGGGCCCCTGAAGATACTTGTTTGGCCCTGCTGTCTATTTAGCCAAGCACCC ACAACAGAGATCAGACCCCAAGTTCTGCTGTTTCAGTTGCCACGCCTACAGTTAGTGTTTCAACT CCTGCTCCTACAGCCACACCTGTGCAAACCGTTCCCCAGCCGCACCCTCAGACGTTACCTCCTGCT GTTCCTCATTCAGTACCTCAGCCAACAACAGCAATACCTGCTTTTTCCACCAGTAATGGTACCTCCG TTTCGTGTTCCCCTTCCTGGCATGCCAATTCCACTTCCAGGTGTATTGCCAGGAATGGCCCCTCCT ATCGTAC_CATGATACATCCCCAGGTTGCTATTGCAGCTTCACCTGCTACCTTAGCTGGAGCAACA GCAGTTTCTGAATGGACTGAATATAAAACAGCAGATGGGAAGACATATTATTATAATAATAATAGAACA $\verb|TTAGAATCAACCTGGGAAAAACCCCCAAGAACTAAAGGAAAAAGAAAAGTTAGAAGAGAGATTAAA$ $\tt GAGCCAATTAAAGAACCCTCTGAAGAGCCTCTGCCAATGGAGACGGAGGAGGAGGATCCTAAAGAA$ GAGCCTATAAAGGAGATAAAGGAGGAGCCCAAAGAAGAGGAGATGACTGAAGAAGAAAAGGCTGCC CAGAAGGCAAAGCCAGTTGCTACTGCTCCTATTCCTGGTACTCCATGGTGTGTCGTTTGGACTGGT ATTGGCAGGCAGATGTTGACAAAATTATTCAGGAGCCCCCTCATAAAAAAGGAATGGAGGAATTG AAGAAACTAAGGCACCCAACTCCGACAATGCTGTCGATCCAAAAGTGGCAATTCTCTATGAGTGCA ATTAAAGAGAACAAGAATTAATGGAAGAAATTAATGAAGATGAGCCTGTTAAAGCAAAAAAACGG AAG

Figure 6 (continued)

>CGI-125

>CGI-74

>CLH-17

>CLK1

Figure 6 (continued)

>DRP-1

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CACCTGTACCAGCGGTCAAAATCAGGAATAAGGTTTTTTGGATTGCAAGGGGTTTCCAGGGGCATG
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TCTTCGCCTTCTAAACACCAGCCCCCACCCATCAGAAACCTCCACCAGTCCAACTTCAGCTTATCA
GGTGCCCAGATAGATGACAACAATGCCAGGCGCCCCCCTGGTGGC
CGCTCCAACATCACCAGCCTCCGT

>EF1A

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GCTGGCTTCACTGCTCAGGTGATTATCCTGAACCATCCAGGCCAAATAAGCGCCGGCTATGCCCCT
GTATTGGATTGCCACACGGCTCACATTGCATGCAAGTTTGCTGAAGCTGAAGGAAAAGATTGATCGC
CGTTCTGGTAAAAAGCTGGAAGATGGCCCTAAATTCTTGAAGTCTGGTGATGCTGCCATTGTTGAT
ATGGTTCCTGGCAAGCCCCATGTGTGTTGAGAGCTTCTCAGACTATCCACCTTTGGGTCGCTTTGCT
GTTCGTGATATGAGACAGACAGTTGCGGTGGGTGTCATCAAAGCAGTGGACAAGAAGGCTGCTGGA
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GCGGCTGGGACCCTGTACACGTATCCTGAAAACTGGAGGGCCTTCAAGGCTCTCATCGCTGCTCAG ACCCCTGAATTTCTCCGCAAATTTCCTGCCGGCAAGGTCCCAGCATTTGAGGGTGATGATGCATTC TGTGTGTTTGAGAGCAACGCCATTGCCTACTATGTGAGCAATGAGGAGCTGCGGGGAAGTACTCCA ACCTGGGTGTTCCCCACCTTGGGCATCATGCACCACAACAACAGGCCACTGAGAATGCAAAGGAG GAAGTGAGGCGAATTCTGGGGCTGCTGGATGCTTACTTGAAGACGAGGACTTTTCTGGTGGGCGAA CGAGTGACATTGGCTGACATCACAGTTGTCTGCACCCTGTTGTGGCTCTATAAGCAGGTTCTAGAG CCTTCTTTCCGCCAGGCCTTTCCCAATACCAACCGCTGGTTCCTCACCTGCATTAACCAGCCCCAG TTCCGGGCTGTCTTGGGCGAAGTGAAACTGTGTGAGAAGATGGCCCAGTTTGATGCTAAAAAGTTT GCAGAGACCCAACCTAAAAAGGACACACCACGGAAAGAGAAGGGTTCACGGGAAGAAGCAGAAG TGTGAGCAGGCGCTGGCTGAGCCCAAGGCCAAGGACCCCTTCGCTCACCTGCCCAAGAGTACC TTTGTGTTGGATGAATTTAAGCGCAAGTACTCCAATGAGGACACACTCTCTGTGGCACTGCCATAT ttctgggagcactttgataaggacggctggtccctgtggtactcagagtatcgcttccctgaagaa CTCACTCAGACCTTCATGAGCTGCAATCTCATCACTGGAATGTTCCAGCGACTGGACAAGCTGAGG AAGAATGCCTTCGCCAGTGTCATCCTTTTTGGAACCAACAATAGCAGCTCCATTTCTGGAGTCTGG GTCTTCCGAGGCCAGGAGCTTGCCTTTCCGCTGAGTCCAGATTGGCAGGTGGACTACGAGTCATAC ACATGGCGGAAACTGGATCCTGGCAGCGAGGAGACCCAGACGCTGGTTCGAGAGTACTTTTCCTGG GAGGGGCCTTCCAGCATGTGGGCAAAGCCCTTCAATCAGGGCAAGATCTTCAAG

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TGGTCCTATGAAGGGCTGAGGCACATGTCTGGGTCTGAGCTGACCCGAGCTGCTGGACCAGGTGGAG
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>G451P1

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AGGACCTGTATGCCCTATATATTTTCTCTGTCCTTGGAGGCTCTGAAATGTTTCCGCATCAGGAAC AATGAGAAGATGCTGAGTGACAGCCACGGCGTGGAGACCATCCGGGACATCCTGCCAGACACCACC CTTGGGGGCCCATCCTTCTTCAAAATCATCACGGCCAAGGCTGTCCTGAAGCTGCAGGCCGGAAAC GCCGAGGAAGCCGCCCTGTGGAGGGATCTGGTCCGCAAAGTCCTGGCATCCTACTTGGAGACAGCC CAGGAGGCGTGACCCTGGGCGGGGGCCTGGATGAAAACTGTCAGGAGGTGGTGAAATTTGGCAGG CGGGAGAATGGCTTCCTGCTGCAGTACCTGGTGGCTATCCCCATGGAGAAAGGCCTTGACTCCCAA GGCTGCTTCTGCGCAGGCTGCTCCCGGCAGATCGGCTTCTCCTTTGTACGACCCAAGCTCTGTGCC TTCTCTGGCCTCTATTACTGTGACATCTGCCACCAAGACGATGCCTCAGTGATTCCGGCCAGGATC CGGGCCCAGCCCTCATCAACCTGCAGATGGTGAACGCGTCTCTGTACGAGCATGTGGAGCGGATG ${\tt CACCTCATTGGGAGGAGCAGCAGCTGAAGCTCCTGGGGGGATTACCTGGGCCTGTGCCGGAGT}$ GGCGCCTGAAGGAGCTCAGCAAGAGGCTCAAGCACAGGAATTATCTCTTGGAATCTCCGCATAGG TTCAGTGTTGCTGACCTCCAACAGATCGCAGACGGGGTGTATGAAGGATTCCTCAAGGCCCTGATT GAATTTGCCTCCCAGCATGTCTACCACTGCGACCTGTGCACCCAGCGCGGCTTCATCTGCCAGATC TGCCAGCACCACGACATCATCTTCCCCTTTGAGTTTGACACCACAGTCAGGTGTGCCGAGTGCAAG ACCGTCTTCCACCAGAGCTGCCAGGCTGTGGTGAAGAAGGGCTGCCCCGGTGTGCCCGCCGGCGC AAGTACCAGGAACAGAACATTTTCGCC

>G45IP3

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Figure 6 (continued)

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>HD1.7

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CCAAGGTTACAGCTCGAGCTCTATAAGGAAATTAAAAAGAATGGTGCCCCTCGGAGTTTGCGTGCT GCCCTGTGGAGGTTTGCTGAGCTGGCTCACCTGGTTCGGCCTCAGAAATGCAGGCCTTACCTGGTG AACCTTCTGCCGTGCCTGACTCGAACAAGCAAGAGACCCGAAGAATCAGTCCAGGAGACCTTGGCT GCAGCTGTTCCCAAAATTATGGCTTCTTTTGGCAATTTTGCAAATGACAATGAAATTAAGGTTTTG TTAAAGCCCTTCATAGCGAACCTGAACTCAAGCTGCGCACATTCGGCGGACAGCGGCTGCATCA GCAGTGAGCATCTGCCAGCACTCAAGAAGGACACAATATTTCTATAGTTGGCTACTAAATGTGCTC TTAGGCTTACTCGTTCCTGTCGAGGATGAACACTCCACTCTGCTGATTCTTGGCGTGCTGCTCACC CTGAGGTATTTGGTGCCCTTGCTGCAGCAGCAGGTCAAGGACACAAGCCTGAAAGGCAGCTTCGGA GTGACAAGGAAAGAAATGGAAGTCTCTCCTTCTGCAGAGCAGCTTGTCCAGGTTTATGAACTGACG TTACATCATACACAGCACCAAGACCACAATGTTGTGACCGGAGCCCTGGAGCTGTTGCAGCAGCTC TTCAGAACGCCTCCACCCGAGCTTCTGCAAACCCTGACCGCAGTCGGGGGCATTGGGCAGCTCACC GCTGCTAAGGAGGAGTCTGGTGGCCGAAGCCGTAGTGGGAGTATTGTGGAACTTATAGCTGGAGGG GGTTCCTCATGCAGCCCTGTCCTTTCAAGAAAACAAAAAGGCAAAGTGCTCTTAGGAGAAGAAGAA GCCTTGGAGGATGACTCTGAATCGAGATCGGATGTCAGCAGCTCTGCCTTAACAGCCTCAGTGAAG GATGAGATCAGTGGAGAGCTGGCTGCTTCTTCAGGGGTTTCCACTCCAGGGTCAGCAGGTCATGAC ATCATCACAGAACAGCCACGGTCACAGCACACTGCAGGCGGACTCAGTGGATCTGGCCAGCTG

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Figure 6 (continued)

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CCCTGCTGCTCTGAGGACACCATCCCTTCCCAAGTTTCAGATTATGATTATTTCTCTGTAAGTGGT CACCAGGAGGCAGATCAGCAGGAGTTCGACAAGTCCTCCACCATTCCAAGAAACAGCGACATCAGC CAGTCCTACCGACGGATGTTCCAAGCCAAGCGTCCAGCCTCAACTGCTGGCCTCCCCACCACCCTG GGACCTGCTATGGTCACTCCAGGGGTTGCAACTATCCGACGGACCCCTTCCACCAAGCCTTCTGTC CGCCGGGGAACCATTGGAGCTGGTCCCATCCCCATCAAGACACCCGTGATCCCTGTCAAGACCCCA ACCGTCCCAGACCTCCCAGGGGTGTTGCCAGCCCCTCCAGATGGGCCAGAAGAGCGGGGGGGAGCAC AGCCCTGAGTCGCCATCTGTGGGTGAGGGCCCCCAAGGTGTCACCAGCATGCCCTCCTCAATGTGG AGCGGCCAAGCTTCCGTTAACCCTCCACTTCCAGGCCCGAAGCCCAGTATCCCTGAGGAGCACAGA CAGGCAATTCCAGAAAGTGAAGCTGAAGACCAGGAACCGGGAACCCCCAAGTGCCACTGTCTCCCCA GGCCAGATTCCAGAGAGTGACCCTGCAGACCTGAGCCCAAGGGATACTCCACAAGGAGAAGACATG CTGAACGCCATCCGAAGGGGCGTGAAACTGAAGAAGACCACGACAAACGATCGCTCAGCCCCTCGC TTTTCT

>HIP15

ATTCACATGGCTCCACCTTATCCAAATCTAAACATGATTGAGACATTCATATGTCAAGTGTGTGAG GAAACCCTTGCACATAGTGTGGATTCCCTTGAGCAGCTGACTGGAATAAGGATGCTTAGACACCTC ACTATGACTATTGACTATCACACACTGATTGCCAACTATATGTCCGGGTTTCTCTCTTATTAACC ACAGCCAATGCGAGAACGAAGTTTCACGTTCTGAAAATGCTATTGAATTTGTCTGAAAATCCTGCT GTGGCAAAAAACTATTCAGTGCCAAAGCTCTTTCAATATTTGTGGGTCTCTTTAACATAGAAGAG **ACAANTGATAATATTCAAATTGTTATTAAAATGTTTCAGAATATCAGTAACATTATAAAAAGTGGA** AAGATGTCCTTAATTGATGATGATTTCAGTCTTGAGCCGCTTATTTCTGCATTTCGTGAATTTGAG GAGTTAGCTAAGCAACTACAAGCCCAAATAGACAACCAAAATGATCCTGAGGTGGGACAACAAAGT

GATGAAGAGGAGAAACCATAGGCAGATGATAAAGGAAGCTTTTGCTGGGGATGATGTCATCAGA GATTTCTTGAAAGAGAAGAGGGAAGCTGTGGAGGCGAGTAAGCCAAAGGACGTGGACCTGACACTA CCTGGCTGGGGCGAGTGGGGTGTGGGGCCTAAAGCCCAGTGCCAAGAAAAGACGCCGGTTTCTC ATTAAAGCCCCTGAGGGTCCTCCAAGAAAAGATAAGAATTTGCCAAATGTGATTATCAATGAGAAG CGCAACATCCACGCAGCTGCTCATCAGGTACGAGTGCTTCCATATCCATTTACCCACCATTTGGCAA TTTGAAAGGACCATCCAGACCCCCATAGGATCCACATGGAACACCCCAGAGGGCTTTCCAAAAGCTG ACTACTCCCAAGGTCGTCACCAAGCCAGGCCATATCATTAACCCCCATAAAAGCAGAAGACGTGGGC TACCGGTCTTCCTCAAGGTCGGACCTGTCTGTCATACAGAGGAATCCAAAACGAATCACCACACGT CACAAAAACAGCTGAAGAAATGCTCTGTAGAT

AGCAAAAATCAAATTAAAGTAGATCTTGTAGATGAGAATTTTACAGAAATTAAGAGGACAAATAGCA ggacctccagacacaccatatgaaggaggaagataccaactagagataaaaataccagaaacatac CCATTTAATCCCCCTAAGGTCCGGTTTATCACTAAAATATGGCATCCTAATATTAGTTCCGTCACA GGGGCTATTTGTTTGGATATCCTGAAAGATCAATGGGCAGCTGCAATGACTCTCCGCACGGTATTA TTGTCATTGCAAGCACTATTGGCAGCTGCAGAGCCAGATGATCCACAGGATGCTGTAGTAGCAAAT $\tt CAGTACAAACAAAATCCCGAAATGTTCAAACAGACAGCTCGACTTTGGGCACATGTGTATGCTGGA$ GCACCAGTTTCTAGTCCAGAATACACCAAAAAAATAGAAAACCTATGTGCTATGGGCTTTGATAGG AATGCAGTAATAGTGGCCTTGTCTTCAAAATCATGGGATGTAGAGACTGCAACAGAATTGCTTCTG AGTAACTG

Figure 6 (continued)

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>HIP5 (bait)

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AATCAGAGCTTTAAGTTTGGAAATCAAAAAGCAGCAGCTATCAGAGATAGTATTGAATTAACAAAG GAAAAAGGTGCAGAAATTCCAAAGACTATTAAAAAACTGAGGTGGTTTGATGAAACTAGCAATATA GAAAACAATGCTGAAAACAGTCATTCACTGAAGAATAAAACAGGAACAACTCAACAGCATTCTCAA CAATTCCACATTCAAAGTGGTGCTGGAAGCAACATAATTAGTGTTTCTACTTGTGCTGTAAATTCT GCTGATACAAGAAGTCCAGGGAGGATTCTATCTCTGAAAATGTTACGACTTTAGGAGGATCTGGA GCAGACCATATGCCTTTGAACTGTTTTATACCTTCAGGTTATAACTTTGCTAAACATGCCTGGCCA . GCCTCAAAAAAAGAAGAAAGTAAAATCCCTGTACATGATGATTCTAAAACTAAGCAAGGTAAGCCA CAAAGAGGTAGAGCAAAAATAATTAGAAAAACCAGGATCTGCAAAAGTCCAATCAGGCTTTATATGT CAGGGAAAATTAATTATACCTTGTCCTCCTCAATCTACATCAAATATTAGAAGTGGTAAAAAAT ATACAAGTGTCTCAGTGTCAACCAGTAACTCCTGAAAATCCTCAAAACATTATTACACATAACTCT TTTAATTCAAAACATGTGCTTCCAACAGAACACAGTTTGAATCAGTGGAATCAGGAAAGTAGTTCT CCACTCTCAAATGCTTGTTCTGACCTAGTCACTGTGATACCATCACTGCCATCATATTGTTCTTCA GAGTGC_AAACTTTCGCAAAAATAAATCATTCAAATGGCACTCAAGCAGTTGCCCGGCAAGATGCG ACATTATATTGCACCCAAAGAAGTCCTGTTTGTGAAGAAAGTTATCCGTCTGTGACTCTAAGAACT GCTGAAGAAGAATCAGTTCCCTTATGGAAAAGAGGTCCTAATGTCCTGCATCAAAATAAGAGGGCT ACAGGGTCTACTGTTATGAGAAAAACGAATTGCTGAAACTAAGCGGAGAAATATTTTAGAGCAG AAAAGACAAAACCCTGGATCTGTAGGACAGAAGTACAGTGAGCAAATTAATAATTTTTGGACAAAGT GTCCTGCTAAGTTCAAGTGAGCCAAAACAAACTACAAGGGGTACTTCTTATATTGAAGAAGTTTCA GATAGTACTTCTGAGTTTTTGATGGCTGAAAACTTAGTGAAAGCATCAGTGCCGGAGGATGAGATT AÁCATCTGCACACTGTCAGCTGAAGAACAGAAGATCCTAGAGTCCCTTAATGATCTCAATGAAGA CTTCTGGAGAGAGAGAGATAGAACCAGCAGCTGCAGAGACAAGAGA

>HMP CAAGAACAAGTTAAAATTGAGTCTCTAGCCAAGAGCTTAGAAGATGCTCTGAGGCAAACTGCAAGT GTCACTCTGCAGGCTATTGCAGCTCAGAATGCTGCGGTCCAGGCTGTCAATGCACACTCCAACATA TTGAAAGCCGCCATGGACAATTCTGAGATTGCAGGCGAGAAGAAATCTGCTCAGTGGCGCACAGTG CAGGGTGCATTGAAGGAACGCAGAAAGGCAGTAGATGAAGCTGCCGATGCCCTTCTCAAAGCCAAA CAAGAGTTAGAGAAGATGAAAAGTGTGATTGAAAATGCAAAGAAAAAAAGAGGTTGCTGGGGGCAAG... CCTCATATAACTGCTGCAGAGGGTAAACTTCACAACATGATAGTTGATCTGGATAATGTGGTCAAA AAGGTCCAAGCAGCTCAGTCTGAGGCTAAGGTTGTATCTCAGTATCATGAGCTGGTGGTCCAAGCT ACTGTTTCAGACTTAGCTGACAAGCTCTCTACTGATGATCTGAACTCCCTCATTGCTCATGCACAT CGTCGTATTGATCAGCTGAACAGAGAGCTGGCAGAACAGAAGGCCACCGAAAAGCAGCACATCACG TTAGCCTTGGAGAAACAAAAGCTGGAAGAAAAGCGGGCATTTGACTCTGCAGTAGCAAAAGCATTA CAACATCACAGAAGTGAAATACAGGCTGAACAGGACAGAAAGATAGAAGAAGTCAGAGATGCCATG GAAAATGAAATGAGAACCCAGCTTCGCCGACAGGCAGCTGCCCACACTGATCACTTGCGAGATGTC CTTAGGGTACAAGAACAGGAATTGAAGTCTGAATTTGAGCAGAACCTGTCTGAGAAACTCTCTGAA CAAGAATTACAATTTCGTCGTCTCAGTCAAGAGCAAGTTGACAACTTTACTCTGGATATAAATACT GCCTATGCCAGACTCAGAGGAATCGAACAGGCTGTTCAGAGCCATGCAGTTGCTGAAGAGGAAGCC AGAAAAGCCCACCAACTCTGGCTTTCAGTGGAGGCATTAAAGTACAGCATGAAGACCTCATCTGCA GAAACACCTACTATCCCGCTGGGTAGTGCGGTTGAGGCCATCAAAGCCAACTGTTCTGATAATGAA TTCACCCAAGCTTTAACCGCAGCTATCCCTCCAGAGTCCCTGACCCGTGGGGTGTACAGTGAAGAG AGAAATAGCTTGTACCAGTACTTCCTCCTACCTACAGTCCCTGCTCCTATTCCCACCTCAGCAA CTGAAGCCGCCCCAGAGCTCTGCCCTGAGGATATAAACACATTTAAATTACTGTCATATGCTTCC TATTGCATTGAGCATGGTGATCTGGAGCTAGCAGCAAAGTTTGTCAATCAGCTGAAGGGGGGAATCC AGACGAGTGGCACAGGACTGGCTGAAGGAAGCCCGAATGACCCTAGAAACGAAACAGATAGTGGAA ATCCTGACAGCATATGCCAGCGCCGTAGGAATAGGAACCACTCAGGTGCAGCCAGAG

>HSPC232 CGGCGGCGAGCGGACGGCTGCATTTACGGGGTCTCCCGGAGGGCCAGAGTCGTGGCTTACAGAAGA GACGAAATGTGGTCTGAGGGACGATATGAATATGAAAGAATTCCGAGAGAACGAGCACCTCCTCGA AGTCATCCCAGTGATGAATCTGGTTATAGATGGACAAGAGACGATCATTCTGCAAGCAGGCAACCT GAATACAGGGACATGAGAGATGGCTTTAGAAGAAAAAGTTTCTACTCTTCCCATTATGCGAGAGAG CGGTCTCCTTATAAAAGGGACAATACTTTTTTCAGAGAATCACCTGTTGGCCGAAAGGATTCTCCA CACAGCAGATCTCCAGTGTCAGTAGCAGAAGCTACTCTCCAGAAAGGAGCAAATCATACTCT TTCCATCAGTCTCAACATAGAAATAAAGAGAGGCCTGTCCAGTCTTTGAAAACATCAAGAGATACT TCACCCTCAAGTGGTTCAGCAGTTTCTTCATCAAAGGTGTTAGACAAACCCAGTAGGCTAACTGAA AAGGAACTTGCTGAGGCTGCAAGCAAGTGGGCTGCTGAAAAGCTAGAGAAATCAGATGAAAGTAAC $\tt CCTGAGTCAAACACACACACGGGATAGAATTATTTGAAGATAGTCAGCTAACCACTCGCTCTAAA$ GCAATAGCATCAAAAACCAAAGAGATTGAACAGGTTTACCGACAAGACTGTGAAAACTTTCGGGATG GTGGTGAAAATGCTGATTGAAAAAGATCCTTCATTAGAAAAGTCTATACAGTTTGCATTGAGGCAG AATTTACATGAAATAGGTGAGCGGTGTGTTGAAGAACTCAAGCATTTCATTGCAGAGTATGATACT TCCACTCAAGATTTTGGAGAGCCTTTT

GGCCGCCGGCGAGCAGTCTGAGCCCGACGATGAGGCCCGGGGACGGGAGCTGAGCGTGGAGGCCTC ATGATGGGGCACCCTGGCATGCATTATGCCCCAATGGGAATGCACCCTATGGGTCAGAGAGCGAAT ATGCCTCCTGTACCTCATGGAATGATGCCCGCAGATGATGCCCCCTATGGGAGGGCCACCAATGGGA CAAATGCCTGGAATGATGTCGTCAGTAATGCCTGGAATGATGATGTCTCATATGTCTCAGGCTTCC ATGCAGCCTGCCTTACCGCCAGGAGTAAATAGTATGGATGTAGCAGCAGGTACAGCATCTGGTGCA AAATCAATGTGGACTGAACATAAATCACCTGATGGAAGGACTTACTACTACAACACTGAAACCAAA $\tt TGGGCCAAACCTAAAGAACTTGAGGATCTTGAAGGATACCAGAATACCATTGTTGCTGGAAGTCTT$ ATTACAAAATCAAACCTGCATGCAATGATCAAAGCTGAAGAAAGCAGTAAGCAAGAAGAGTGCACC ACAACATCAACAGCTCCAGTCCCTACAACAGAAATTCCGACCACAATGAGCACCATGGCTGCTGCC GAAGCAGCAGCTGCTGTTGTTGCAGCAGCAGCAGCAGCAGCAGCAGCAGCAGCAGCAATGCT AATGCTTCCACTTCTGCTTCTAATACTGTCAGTGGAACTGTTCCAGTTGTTCCTGAGCCTGAAGTT ACTTCCATTGTTGCTACTGTTGTAGATAATGAGAATACAGTAACTATTTCAACTGAGGAACAAGCA CAACTTACTAGTACCCCTGCTATTCAGGATCAAAGTGTGGAAGTATCCAGTAATACTGGAGAAGAA ACATCTAAGCAAGAAACTGTAGCTGATTTTACTCCCAAAAAAGAAGAGGAGGAGGAGCCAACCAGCA AAGCGGGTACCATCGAATGCTTCATGGGAGCAGGCTATGAAAATGATTATTAATGATCCACGATAC AGTGCTTTGGCAAAGTTAAGTGAAAAAAGCAAGCCTTTAATGCCTATAAAGTCCAGGCAAAAAA

AAAGAAAAAAAAAAAAAAAAAAAA

CACGCCGCTTCGCCGAGGCCGAGTGCCTGGCCGAGAGCCACCAGCACCTCTCCAAGGAGTCGCTG GCGGGGAACAAGCCGGCCAACGCCGTCCTGCACAAGGTTCTGAACCAGCTGGAGGAGTTGCTGAGC GACATGAAGGCGGACGTGACCCGCCTGCCAGCCACGCTGTCCCGAATACCCCCCATCGCAGCCCGC CTTCAGATGTCCGAGCGCAGCATCCTCAGCCGGCTGGCCAGCAAGGGCACGGAGCCTCACCCCACA GTAGGGGCCCTGGCCGCCGCGGGCGCCAATTACAGCCAGATGCCTGCAGGGTCCTTCATCACAGCC GGGCTGGATCGGAAGGAGCCCCGAGCCGGGGAGGTGATCTGTATAGACGAC

>IKAP
CTCAAAGAAGGCAGTCCGCTGGAGGACCTGGCCCTCCTGGAGGCACTGAGTGAAGTGGTGCAGAAC
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GACTTTCTCATTCTCCCTGGGTACATCGACTTCACTGCAGACCAGGTGGACCTGACTTCTGCTCTG ACCAAGAAAATCACTCTTAAGACCCCACTGGTTTCCTCTCCCATGGACACAGTCACAGAGGCTGGG ATGGCCATAGCAATGGCGCTTACAGGCGGTATTGGCTTCATCCACCACAACTGTACACCTGAATTC CAGGCCAATGAAGTTCGGAAAGTGAAGAATATGAACAGGGATTCATCACAGACCCTGTGGTCCTC AGCCCCAACCATCCCCTGCCCCATCTTTTTCACCCCAACGCCCCATCCTTTCTCCCCCAACCCA ATCACAGACACAGGCCGGATGGGGAGCCGCTTGGTGGGCATCATCTCCTCCAGGGACATTGATTTT CTCAAAGAGGGGGAACATGACTGTTTCTTGGAAGAGATAATGACAAAGAGGGAAGACTTGGTGGTA GCCCCTGCAGGCATCACACTGAAGGAGGCAAATGAAATTCTGCAGCGCAGCAAGAAGGGAAAGTTG CCCATTGTAAATGAAGATGATGAGCTTGTGGCCATCATTGCCCGGACAGACCTGAAGAAGAATCGG GACTACCCACTAGCCTCCAAAGATGCCAAGAAACAGCTGCTGTGTGGGGGCAGCCATTGGCACTCAT CAGGATGACAAGTATAGGCTGGACTTGCTCGCCCAGGCTGGTGTGGATGTAGTGGTTTTTGGACTCT TCCCAGGGAAATTCCATCTTCCAGATCAATATGATCAAGTACATCAAAGACAAATACCCTAATCTC CAAGTCATTGGAGGCAATGTGGTCACTGCTGCCCAGGCCAAGAACCTCATTGATGCAGGTGTGGAT GCCCTGCGGGTGGGCATGGGAAGTGGCTCCATCTGCATTACGCAGGAAGTGCTGGCCTGTGGGCGG CCCCAAGCAACAGCAGTGTACAAGGTGTCAGAGTATGCACGGCGCTTTTGGTGTTCCCGGTCATTGCT GATGGAGGAATCCAAAATGTGGGTCATATTGCGAAAGCCTTGGCCCCTTGGGGCCCTCCACAGTCATG ATGGGCTCTCTCCTGGCTGCCACCACTGAGGCCCCTGGTGAATACTTCTTTTCCGATGGGATCCGG CTAAAGAAATATCGCCGCTATGGGTTCTCTCGATGCCATGGACAAGCACCTCAGCAGCCAGAACAGA TATTTCAGTGAAGCTGACAAAATCAAAGTGGCCCAGGGAGTGTCTGGTGCTGTGCAGGACAAAGGG TCAATCCACAAATTTGTCCCTTACCTGATTGCTGGCATCCAACACTCATGCCAGGACATTGGTGCC AAGAGCTTGACCCAAGTCCGAGCCATGATGTACTCTGGGGGAGCTTAAGTTTGAGAAGAGAACGTCC TCAGCCCAGGTGGAAGGTGGCGTCCATAGCCTCCATTCGTATGAGAAGCGGCTTTTC

>KPNA2 GCTTGGGCACTCACTAACATTGCTTCTGGGACATCAGAACAAACCAAGGCTGTGGTAGATGGAGGT GCCATCCCAGCATTCATTTCTCTGTTGGCATCTCCCCATGCTCACATCAGTGAACAAGCTGTCTGG GCTCTAGGAAACATTGCAGGTGATGGCTCAGTGTTCCGAGACTTGGTTATTAAGTACGGTGCAGTT GACCCACTGTTGGCTCTCCTTGCAGTTCCTGATATGTCATCTTTAGCATGTGGCTACTTACGTAAT CTTACCTGGACACTTTCTAATCTTTGCCGCAACAAGAATCCTGCACCCCCGATAGATGCTGTTGAG ${\tt CAGATTCTTCCTACCTTAGTTCGGCTCCTGCATCATGATGATCCAGAAGTGTTAGCAGATACCTGC}$ TGGGCTATTTCCTACCTTACTGATGGTCCAAATGAACGAATTGGCATGGTGGTGAAAACAGGAGTT GTGCCCCAACTTGTGAAGCTTCTAGGAGCTTCTGAATTGCCAATTGTGACTCCTGCCCTAAGAGCC ${\tt AACATCACAGCCGGCCGGCCAGGACCAGATACAGCAAGTTGTGAATCATGGATTAGTCCCATTCCTT}$ GTCAGTGTTCTCTCTAAGGCAGATTTTAAGACACAAAAGGAAGCTGTGTGGGCCCGTGACCAACTAT ACCAGTGGTGGAACAGTTGAACAGATTGTGTACCTTGTTCACTGTGGCATAATAGAACCGTTGATG · AACCTCTTAACTGCAAAAGATACCAAGATTATTCTGGTTATCCTGGATGCCATTTCAAATATCTTT CAGGCTGCTGAGAAACTAGGTGAAACTGAGAAACTTAGTATAATGATTGAAGAATGTGGAGGCTTA GACAAAATTGAAGCTCTACAAAACCATGAAAATGAGTCTGTGTATAAGGCTTCGTTAAGCTTAATT GAGAAGTATTTCTCTGTAGAGGAAGAGGGAAGATCAAAACGTTGTACCAGAAACTACCTCTGAAGGC TACACTTTCCAAGTTCAGGATGGGGCTCCTGGGACCTTTAACTTT

>KPNBJ GACGAGGTGATGCAGCTGCTTCTGGAAAATTTGGGGAATGAGAACGTCCACAGGTCTGTGAAGCCG CAGATTCTGTCAGTGTTTGGTGATATTGCCCTTGCTATTGGAGGAGAGTTTAAAAAATACTTAGAG GTTGTATTQAATACTCTTCAGCAGGCCTCCCAAGCCCAGGTGGACAAGTCAGACTATGACATGGTG GATTATCTGAATGAGCTAAGGGAAAGCTGCTTGGAAGCCTATACTGGAATCGTCCAGGGATTAAAG GGGGATCAGGAGAACGTACACCCGGATGTGATGCTGGTACAACCCAGAGTAGAATTTATTCTGTCT TTCATTGACCACATTGCTGGAGATGAGGATCACACAGATGGAGTAGTAGCTTGTGCTGCTGGACTA ATAGGGGACTTATGTACAGCATTTGGGAAGGATGTACTGAAATTAGTAGAAGCTAGGCCAATGATC CATGAATTGTTAACTGAAGGGCGGAGATCGAAGACTAACAAAGCAAAAACCCTTGCTACATGGGCA ACAAAAGAACTGAGGAAACTGAAGAACCAAGCT

>Ku70

AAGACCCGGACCTTTAATACAAGTACAGGCGGTTTGCTTCTGCCTAGCGATACCAAGAGGTCTCAG ATCTATGGGAGTCGTCAGATTATACTGGAGAAAGAGGGAAACAGAAGAGCTAAAACGGTTTGATGAT $\tt CCAGGTTTGATGCTCATGGGTTTCAAGCCGTTGGTACTGCTGAAGAAACACCATTACCTGAGGCCCC$ TCCCTGTTCGTGTACCCAGAGGAGTCGCTGGTGATTGGGAGCTCAACCCTGTTCAGTGCTCTGCTC ATCAAGTGTCTGGAGAAGGAGGTTGCAGCATTGTGCAGATACACACCCCGCAGGAACATCCCTCCT TATTTTGTGGCTTTGGTGCCACAGGAAGAGAGTTGGATGACCAGAAAATTCAGGTGACTCCTCCA GGCTTCCAGCTGGTCTTTTTACCCTTTGCTGATGATAAAAGGAAGATGCCCTTTACTGAAAAAATC ATGGCAACTCCAGAGCAGGTGGGCAAGATGAAGGCTATCGTTGAGAAGCTTCGCTTCACATACAGA AGTGACAGCTTTGAGAACCCCGTGCTGCAGCAGCACTTCAGGAACCTGGAGGCCTTGGCAT TTGATGGAGCCGGAACAAGCAGTGGACCTGACATTGCCCAAGGTTGAAGCAATGAATAAAAGACTG GGCTCCTTGGTGGATGAGTTTAAGGAGCTTGTTTACCCACCAGATTACAATCCTGAAGGGAAAGTT ACCAAGAGAAAACACGATAATGAAGGTTCTGGAAGCAAAAGGCCCAAGGTGGAGTATTCAGAAGAG GAGCTGAAGACCCACATCAGCAAGGGTACGCTGGGCAAGTTCACTGTGCCCCATGCTGAAAGAGGCC TGCCGGGCTTACGGGCTGAAGAGTGGGCTGAAGAAGCAGGAGCTGCTGGAAGCCCTCACCAAGCAC TTCCAGGAC

>LUC7B1

GTCGACGCGGTCGCGGTCGACGCGGCCGCGGTTTCTGCAAAGGCAGAAAAAGTACATGAGTTAAAT GAAAAAATAGGAAAACTCCTTGCTAAAGCCGAACAGCTAGGGGCTGAAGGTAATGTGGATGAATCC CAGAAGATTCTTATGGAAGTGGAAAAAGTTCGTGCGAAGAAAAAAGAAGCTGAGGAAGAATACAGA AATTCCATGCCTGCATCCAGTTTTCAGCAGCAAAAGCTGCGTGTCTGCGAGGTCTGTTCAGCCTAC CTTGGTCTCCATGACAATGACCGTCGCCTGGCAGACCACTTCGGTGGCAAGTTACACTTGGGGTTC ATTCAGATCCGAGAGAAGCTTGATCAGTTGAGGAAAACTGTCGCTGAAAAGCAGGAGAAGAGAAAAT CAGGATCGCTTGAGGAGGAGAGAGGGAGGGAGGGAGGAGCGTCTGAGCAGGAGGTCGGGATCA AGAACCAGAGATCGCAGGAGGTCACGCTCCCGGGATCGGCGTCGGAGGCGGTCAAGATCTACCTCC CGAGAGCGACGGAAATTGTCCCGGTCCCGGTCCCGAGATAGACATCGGCGCCACCGCAGCCGTTCC CGGAGCCACAGCCGGGACATCGTCGGGGCTTCCCGGGACCGAAGTGCGAAATACAAGTTCTCCAGA CTTCACAGCCTCCAACGGGAAGATGGCTTCACGGAGGTCAGAAGAAGGAGGCCCGGCGAGATC

>MAGEHI

GCATCCTTCCCTAGGACTGCTGTAAGCTTTGAGCCTCTAGCAGGAGACATGCCTCGGGGACGAAAG AGTCGGCGCCGTAATGCGAGAGCCGCAGAAGAGAACCGCAACAATCGCAAAATCCAGGCCTCA GAGGCCTCCGAGACCCCTATGGCCGCCTCTGTGGTAGCGAGCACCCCCGAAGACGACCTGAGCGGC CCCGAGGAAGACCCGAGCACTCCAGAGGAGGCCTCTACCACCCCTGAAGAAGCCTCGAGCACTGCC CAAGCACAAAAGCCTTCAGTGCCCCGGAGCAATTTTCAGGGCACCAAGAAAAGTCTCCTGATGTCT ATATTAGCGCTCATCTTCATCATGGGCAACAGCGCCCAAGGAAGCTCTGGTCTGGAAAGTGCTGGGG AAGTTAGGAATGCAGCCTGGACGTCAGCACAGCATCTTTGGAGATCCGAAGAAGATCGTCACAGAA GAGTTTGTGCGCAGAGGGTACCTGATTTATAAACCGGTGCCCCGTAGCAGTCCGGTGGAGTATGAG TTCTTCTGGGGGCCCCGAGCACACGTGGAATCGAGCAAACTGAAAGTCATGCATTTTGTGGCAAGG ${\tt GTTCGTAACCGATGCTCTAAAGACTGGCCTTGTAATTATGACTGGGATTCGGACGATGATGCAGAG}$ GTTGAGGCTATCCTCAATTCAGGTGCTAGGGGTTATTCCGCCCCT

>MAP11c3

>mHAP1

CCGAAAGAGCAGGTGCAGAGCGGTGCGGGAGACGGGACAGGGTCGGGGGACCCAGCAGCAGCACC CCCACGACCCAGCCTGCAGTTGGTCCCGCTCCGGAGCCCTCGGCGGAGCCCAAACCTGCTCCAGCG CAGGGAACCGGGTCCGGACAAAATCAGGATCCCGAACCAAGACAGGAAGCTTTTGTCGGTCCATG ATCATTGGTGATTCGGACGCACCATGGACCCGCTACGTATTCCAGGGGCCTTACGGTCCCCGGGCC ACTGGCCTGGGCACTGGAAAGGCCGAGGGAATCTGGAAGACAGGAGGCGGCGTACATCGGCCGGAGG CCCGGCGTGTCCGGCCCTGAGCGTGCGGCGTTTATTCGAGAGCTGCAGGAAGCGTTGTGTCCTAAT CCACCACCCACGAAGAAGATCACCGAAGATGATGTCAAAGTGATGTTGTATTTGCTGGAAGAGAAA GAACGGGACCTGAACAGAGCCGCCCGGATCGGCCAGTCCCTGGTGAAACAGAACAGTGTCTTGATG GAGGAGAATAATAAGCTGGAAACCATGCTGGGCTCAGCCAGGGAGGAGATTTTACATCTCCGGAAG CAGGTGAACCTGCGAGATGACCTTCTTCAGCTCTACTCAGACTCTGATGACGATGATGATGAGGAA GACGAGGAAGACGAGGAGGGGCGAAGAGGAGGAACGAGAAGGACAGAGGGGATCAAGACCAGCAG CACGACCACCCCTATGGTGCCCCCAAGCCACACCCTAAGGCTGAGACAGCGCACCGCTGCCCCACAG CTGGAAACCCTGCAGCAGAAGCTCAGGCTTCTGGAGGAAGAGAACGACCACCTGCGAGAGGAGGCC TCCCACCTTGACAACCTGGAGGACGAAGAGCAGATGCTCATTCTGGAATGTGTGGAGCAGTTCTCT GAAGCCAGCCAGCAGATGGCAGAGCTATCGGAAGTGCTGGTGTTGAGGCTGGAAGGCTATGAGAGG CAGCAGAAAGAGATCACTCAGCTGCAGGCCGAGATCACCAAGCTACAACAGCGTTGTCAGTCTTAT GGGGCCCAGACGGAGAAACTGCAGCAGATGCTGGCCTCAGAGAAGGGGGATCCACTCGGAGAGCCCTG CGAGCTGGCTCCTACATGCAGGATTATGGGAGCAGGCCTCGTGACCGCCAGGAGGATGGGAAGAGT CATCGCCAGCGCTCCTCCATGCCCGCAGGCTCTGTCACCCACTATGGATACAGTGTGCCTCTGGAT GCACTTCCAAGTTTCCCAGAGACACTGGCTGAGGAGCTCCGAACATCTCTGAGGAAGTTCATCACT GCGATGCCACCCCCACCGGCTCA

>mp53

Figure 6 (continued)

>NAG4 CGAGACCGGGTGGAGAATGAGGCAGAAAAAGATCTCCAGTGTCACGCCCCTGTGAGATTAGACTTG CCTCCTGAGÁAGCCTCTCACAAGCTCTTTAGCCAAACAAGAAGAAGTAGAACAGACACCCCCTTCAA GTGACTGATTTTATTGCTCCTGGCTACTCCATGATCATTAAACACCCAATGGATTTTAGTACCATG AAAGAAAAGATCAAGAACAATGACTATCAGTCCATAGAAGAACTAAAGGATAACTTCAAACTAATG TGTACTAATGCCATGATTTACAATAAACCAGAGACCATTTATTATAAAGCTGCAAAGAAGCTGTTG CACTCAGGAATGAAAATTCTTAGCCAGGAAAGAATTCAGAGCCTGAAGCAGAGCATAGACTTCATG GCTGACTTGCAGAAAACTCGAAAGCAGAAAGATGGAACAGACACCTCACAGAGTGGGGAGGACGGA ggctgctggcagagagagagagagactctggagatgccgaagcacacgccttcaagagtcccagc AAAGAAAATAAAAAGAAAGACAAAGATATGCTTGAAGATAAGTTTAAAAGCAATAATTTAGAGAGA GAGCAGGAGCAGCTTGACCGCATCGTGAAGGAATCTGGAGGAAAGCTGACCAGGCGGCTTGTGAAC agtcagtgcgaatttgaaagaaga<u>aaaccagatggaacaacg</u>aegttggaggttctccatccttg GATCCCATTGTÄGGAGAGCCAGGCTACTGCCCTGTGAGACTGGGAÄTGÄCAACTGGAAGACTTCAG TCTGGAGTGAATACTTTGCAGGGGTTCAAAGAGGATAAAAGGAACAAAGTCACTCCAGTGTTATAT TTCAATTATGGGCCCTACAGTTCTTATGCACCGCATTATGACTCCACATTTGCAAATATCAGCAAG CATGATTCTGÄTTTÄATCTATTCAACCTATGGGGAAGACTCTGÄTCTTCCAAGTGATTTCAGCATC CATGAGTTTTTGGCCACGTGCCAAGATTATCCGTATGTCATGGCAGATAGTTTACTGGATGTTTTA ACAAAAGGAGGCATTCCAGGACCCTACAAGAGATGGAGATGTCATTGCCTGAAGATGAAGGCCAT ACTAGGACACTTGACACAGCAAAAGAAATGGAGATTACAGAAGTAGAGCCACCAGGGCGTTTGGAC TCCAGTACTCAAGACAGGCTCATAGCGCTGAAAGCAGTAACAAATTTTGGCGTTCCAGTTGAAGTT TTTGACTCTGAAGAAGCTGAAATATTCCAGAAGAAACTTGATGAGACCACCAGATTGCTCAGGGAA CTCCAGGAAGCCCAGAATGAACGTTTGAGCACCAGACCCCCTCCGAACATGATCTGTCTCTTGGGT CCCTCATACAGAGAAATGCATCTTGCTGAACAAGTGACCAATAATCTTAAAGAACTTGCACAGCAA CCCGTCATGGAAAACAACTTTGTGGATTTGACAGAAGACACTGAAGAACCTAAAAAAGACGGATGTT GCTGAGTGTGGACCTGGTGGAAGT

>NEFL CTCTCTCCCCTGTCCTCTCTCCGGGCTCCCACCGCCGCGGGGGCCGGGGAGCCACCGGCCGCC ACCATGAGTTCCTTCAGCTACGAGCCGTACTACTCGACCTCCTACAAGCGGCGCTACGTGGAGACG CCCCGGGTGCACATCTCCAGCGTGCGCAGCGCTACAGCACCGCACGCTCAGCTTACTCCAGCTAC TCGGCGCCGGTGTCTTCCTCGCTGTCCGTGCGCCGCAGCTACTCCTCCAGCTCTGGATCGTTGATG CCCAGTCTGGAGAACCTCGACCTGAGCCAGGTAGCCGCCATCAGCAACGACCTCAAGTCCATCCGC acecaegagaagaccccacctccaegacctcaateacctccaccttcaccttcaccttcacctccaccttcacc GAGCTGGAGCAGCAGAACAAGGTCCTGGAAGCCGAGCTGCTGGTGCTGCGCCAGAAGCACTCCGAG CCATCCCGCTTCCGGGCGCTGTACCAGCAGGAGATCCGCGACCTGCGCCTGGCGGCGGAAGATGCC ACCAACCAGAAGCAGGCGCTCCAGGGCGAGGCGCAAGGGCTGGAGGAGACCCTGCGCAACCTGCAG GCGCGCTATGAAGAGGAGGTGCTGAGCCGCGAGGACGCCGAGGGCCGGCTGATGGAAGCGCGCAAA GGCGCCCACGACGCGCCGCCCCCCCCCCCCCCAGCTCGACAAGCGCATCGACAGCTTGATGGACGAA ATCTCTTTTCTGAAGAAAGTGCACGAAGAGAGAGATCGCCGAACTGCAGGCGCAGATCCAGTACGCG CAGATCTCCGTGGAGATGGACGTGACCAAGCCCGACCTTTCCGCCGCGCTCAAGGACATCCGCGCG CAGTACGAGAAGCTGGCCGCCAAGAACATGCAGAACGCTGAGGAATGGTTCAAGAGCCGCTTCACC GTGCTGACCGAGAGCGCCGCCAAGAACACCGACGCCGTGCGCGCCCAAGGACGAGGTGTCCGAG AGCCGTCGTCTGCTCAAGGCCAAGACCCTGGAAATCGAAGCATGCCGGGGCATGAATCAAGCGCTG GAGAAGCAGCTGCAGGAGCTGGAGGACAAGCAGAACGCCGACATCAGCGCTATGCAGGACACGATC AACAAATTAGAAAATGAATTGAGGACCACAAAGAGTGAAATGGCACGATACCTAAAAGAATACCAA GACCTCCTCAACGTGAAGATGGCTTTGGATATTGAGATTGCAGCTTACAGGAAACTCTTGGAAGGC GAGGAGACCCGACTCAGTTTCACCAGCGTGGGAAGCATAACCAGTGGCTACTCCCAGAGCTCCCAG GTCTTTGGCCGATCTGCCTACGGCGGTTTACAGACCAGCTCCTATCTGATGTCCACCCGCTCCTTC CCGTCCTACTACACCAGCCATGTCCAAGAGGAGCAGATCGAAGTGGAGGÀAACCATTGAGGCTGCC GAAGAGGCCGAGGAAGAGGCAGCTGAAGAGGAAGAAGCTGCCAAGGAAGAGTCTGAAGAAGCA AAAGAAGAAGAAGAAGGAGGTGAAGGAGAAGGAGGAAACCAAAGAAGCTGAAGAGGAGGAG

 ${\tt ATGGAGGAGCCGCAGTCAGATCCTAGCGTCGAGCCCCCTCTGAGTCAGGAAACATTTTCAGACCTA}$ TGGAAACTACTTCCTGAAAACAACGTTCTGTCCCCCTTGCCGTCCCAAGCAATGGATGATTTGATG CTGTCCCCGGACGATATTGAACAATGGTTCACTGAAGACCCAGGTCCAGATGAAGCTCCCAGAATG CCAGAGGCTGCTCCCCCGTGGCCCCTGCACCAGCAGCTCCTACACCGGCGGCCCCTGCACCAGCC CCCTCCTGGCCCCTGTCATCTTCTGTCCCTTCCCAGAAAACCTACCAGGGCAGCTACGGTTTCCGT CTGGGCTTCTTGCATTCTGGGACAGCCAAGTCTGTGACTTGCACGTACTCCCCTGCCCTCAACAAG ACCCGCGTCCGCGCCATGGCCATCTACAAGCAGTCACAGCACATGACGGAGGTTGTGAGGCGCTGC CCCCACCATGAGCGCTGCTCAGATAGCGATGGTCTGGCCCCTCCTCAGCATCTTATCCGAGTGGAA GGAAATTTGCGTGTGGAGTATTTGGATGACAGAAACACTTTTCGACATAGTGTGGTGGTGCCCTAT GAGCCGCCTGAGGTTGGCTCTGACTGTACCACCATCCACTACAACTACATGTGTAACAGTTCCTGC ATGGGCGGCATGAACCGGAGGCCCATCCTCACCATCATCACACTGGAAGACTCCAGTGGTAATCTA $\tt CTGGGACGGAACAGCTTTGAGGTGCGTGTTTGTGCCTGTCCTGGGAGAGACCGGCGCACAGAGGAAA$ GAGAATCTCCGCAAGAAAGGGGAGCCTCACCACGAGCTGCCCCCAGGGAGCACTAAGCGAGCACTG CAGATCCGTGGGCGTGAGCGCTTCGAGATGTTCCGAGAGCTGAATCAGGCCTTGGAACTCAAGGAT GCCCAGGCTGGGAAGGAGCCAGGGGGGAGCAGGGCTCACTCCAGCCACCTGAAGTCCAAAAAGGGT CAGTCTACCTCCCGCCATAAAAACTCATGTTCAAGACAGAAGGGCCTGACTCAGAC

Figure 6 (continued)

>PFN2
GCTCCTCGCCGTCCGCGCTGCAGTGCGAAGGGCTCGAAGATGGCCGGTTGGCAGAGCTACGTGGAT
AACCTGATGTGCGATGGCTGCCAGGAGGCCGCCATTGTCGGCTACTGCGACGCCAAATACGTC
TGGGCAGCCACGGCCGGGGGCGTCTTTCAGAGCATTACGCCAATAGAAATAGATATGATTGTAGGA
AAAGACCGGGAAGGTTTCTTTACCAACGGTTTGACTCTTTGGCGCGAAGAAATGCTCAGTGATCAGA
CATAGTGTATACGTCGATGGTGACTGCACAATGGACATCCGGACAAAAGAAGTCAAGGTGGGCAAGACA
ACATACAATGTGGCTGTCGGCAGAGCTGGTAGAGTCTTTGGTCTTTTGTAATGGCAAAAGAAGGGGTC
CATGGAGGCGGATTGAATAAGAAGGCATACTCAATGGCAAAATACTTGAGAGACTCTGGGTTC
>PIASy(bait)

CTGGTGGAGGCCAAAAACATGGTGATGAGTTTTCGAGTCTCCGACCTTCAGATGCTCCTGGGTTTC GTGGGCCGGAGTAAGAGTGGACTGAAGCACGAGCTCGTCACCAGGGCCCTCCAGCTGGTGCAGTTT GACTGTAGCCCTGAGCTGTTCAAGAAGATCAAGGAGCTGTACGAGACCCGCTACGCCAAGAAGAAC TCGGAGCCTGCCCCACAGCCGCCCCGGCCCCTGGACCCCTGACCATGCACTCCACCTACGACCGG GCCGGCGCTGTGCCCAGGACTCCGCTGGCAGGCCCCAATATTGACTACCCCGTGCTCTACGGAAAG TACTTAAACGGACTGGGACGGTTGCCCGCCAAGACCCTCAAGCCAGAAGTCCGCCTGGTGAAGCTG $\tt CCGTTCTTTAATATGCTGGATGAGCTGCTGAAGCCCCACCGAATTAGTCCCCACAGAACAACGAGAAG$ CTTCAGGAGAGCCCGTGCATCTTCGCATTGACGCCAAGACAGGTGGAGTTGATCCGGAACTCCAGG GAACTGCAGCCCGGAGTTAAAGCCGTGCAGGTCGTCCTGAGAATCTGTTACTCAGACACCAGCTGC CCTCAGGAGGACCAGTACCCGCCCAACATCGCTGTGAAGGTCAACCACAGCTACTGCTCCGTCCCG GGCTACTACCCCTCCAATAAGCCCGGGGTGGAGCCCAAGAGGCCGTGCCGCCCCATCAACCTCACT CACCTCATGTACCTGTCCTCGGCCACCAACCGCATCACTGTCACCTGGGGGAACTACGGCAAGAGC TACTCGGTGGCCCTGTACCTGGTGCGGCAGCTGACCTCATCGGAGCTGCTGCAGAGGCTGAAGACC ATTGGGGTAAAGCACCCGGAGCTGTGCAAGGCACTGGTCAAGGAGAAGCTGCGCCTTGATCCTGAC AGCGAGATCGCCACCACCGGTGTGCGGGTGTCCCTCATCTGTCCGCTGGTGAAGATGCGGCTCTCC GTGCCCTGCCGGCAGAGACCTGCGCCCACCTGCAGTGCTTCGACGCCGTCTTCTACCTGCAGATG ATCGACGGGCTCCTCTCGAAGATCCTGAGCGAGTGTGAGGACGCCGACGAGATCGAGTACCTGGTG GACGGCTCGTGGTGCCCGATCCGCGCCGAAAAGGAGCGCAGCTGCAGCCCGCAGGGCGCCATCCTC GGCAGCACGGGTGGCGGCGCCCGGTGGGCAGCATGGAGAATGGGAAGCCGGGCGCCGATGTGGTG GACCTCACGCTGGACAGCTCATCGTCCTCGGAGGATGAGGAGGAGGAGGAAGAGAGGAGGAAGAC GAGGACGAAGAGGGCCCCGGCCCAAGCGCCCGCTGCCCCTTCCAGAAGGGCCTGGTGCCGGCCTGC

Figure 6 (continued)

WO 2004/113566 PCT/EP2004/006617

>PIASy(prey) CTGGTGGAGGCCAAAAACATGGTGATGAGTTTTCGAGTCTCCGACCTTCAGATGCTCCTGGGTTTC GTGGGCCGGAGTAAGAGTGGACTGAAGCACGAGCTCGTCACCAGGGCCCTCCAGCTGGTGCAGTTT GACTGTAGCCCTGAGCTGTTCAAGAAGATCAAGGAGCTGTACGAGACCCGCTACGCCAAGAAGAAC TCGGAGCCTGCCCCACAGCCGCACCGGCCCCTGGACCCCTGACCATGCACTCCACCTACGACCGG Gecccoeffctgecaggactccgcaggccccaatattgactaccccgtgctctacggaaag TACTTAĀĀCGGĀCTGGGACGGTTGCCCGCCAAGACCCTCAAGCCĀGĀAGTCCGCCTGGTGAAGCTĠ CCGTTCTTTAATATGCTGGATGAGCTGCTGAAGCCCACCGAATTAGTCCCACAGAACAACGAGAAG CTTCAGGAGAGCCCGTGCATCTTCGCATTGACGCCAAGACAGGTGGAGTTGATCCGGAACTCCAGG GAACTGCAGCCCGGAGTTAAAGCCGTGCAGGTCGTCCTGAGAATCTGTTACTCAGACACCAGCTGC CCTCAGGAGGACCAGTACCCGCCCAACATCGCTGTGAAGGTCAACCACAGCTACTGCTCCGTCCCG GGCTACTACCCCTCCAATAAGCCCGGGGTGGAGCCCAAGAGGCCGTGCCGCCCCATCAACCTCACT CACCTCATGTACCTGTCCTCGGCCACCAACCGCATCACTGTCACCTGGGGGAACTACGGCAAGAGC TACTCGGTGGCCCTGTACCTGGTGCGGCAGCTGACCTCATCGGAGCTGCTGCAGAGGCTGAAGACC ATTGGGGTAAAGCACCCGGAGCTGTGCAAGGCACTGGTCAAGGAGAAGCTGCGCCTTGATCCTGAC AGCGAGATCGCCACCACCGGTGTGCGGGTGTCCCTCATCTGTCCGCTGGTGAAGATGCGGCTCTCC GTGCCCTGCGGGCAGAGACCTGCGCCCACCTGCAGTGCTTCGACGCCGTCTTCTACCTGCAGATG ATCGACGGGCTCCTCTCGAAGATCCTGAGCGAGTGTGAGGACGCCGACGAGATCGAGTACCTGGTG GACGGCTCGTGGTGCCCGATCCGCGGAAAAGGAGCGCAGCTGCAGCCCGCAGGGGCGCCATCCTC GCCAGCACGGGTGGCGGCCCGGTGGGCAGCATGGAGAATGGGAAGCCGGGCGCCCCATGTGGTG GACCTCACGCTGGACAGCTCATCGTCCTCGGAGGATGAGGAGGAGGAGGAAGAGGAGGAAGAC GAGGACGAAGAGGGCCCCGGCCCAAGCGCCGCTGCCCCTTCCAGAAGGGCCTGGTGCCGGCCTGC

GGGGAGATAATCGAGGGCTGCCGCCTACCCGTGCTGCGGGGGAACCAGGACAACGAAGATGAGTGG CCCCTGGCCGAGATCCTGAGCGTGAAGGACATCAGTGGCCGGAAGCTTTTCTACGTCCATTACATT GACTTCAACAAACGTCTGGATGAATGGGTGACGCATGAGCGGCTGGACCTAAAGAAGATCCAGTTC CCCAAGAAAGAGGCCAAGACCCCCACTAAGAACGGACTTCCTGGGTCCCGTCCTGGCTCTCCAGAG AGAGAGGTGAAACGGAAGGTGGAGGTGGTTTCACCAGCAACTCCAGTGCCCAGCGAGACAGCCCCG AAATCGAATTGTTTGGGCACTGATGAGGACTCCCAGGACAGCTCTGATGGAATACCGTCAGCACCA CGCATGACTGGCAGCCTGGTGTCTCATCGAAGCCACGACGACATCGTCACCCGGATGAAGAACATT GAGTGCATTGAGCTGGGCCGCCACCGCCTCAAGCCGTGGTACTTCTCCCCGTACCCACAGGAACTC ACCACATTGCCTGTCCTCTACCTGTGCGAGTTCTGCCTCAAGTACGGCCGTAGTCTCAAGTGTCTT CAGCGTCATTTGACCAAGTGTGACCTACGACATCCTCCAGGCAATGAGATTTACCGCAAGGGCACC ATCTCCTTCTTTGAGATTGATGGACGTAAGAACAAGAGTTATTCCCAGAACCTGTGTCTTTTGGCC AAGTGTTTCCTTGACCATAAGACACTGTACTATGACACAGACCCTTTCCTCTTCTACGTCATGACA TACAATGTGGCCTGCATCCTAACCCTGCCTCCCTACCAGCGCCGGGGCTACGGCAAGCTGCTGATC GAGTTCAGCTATGAACTCTCCAAAGTGGAAGGGAAAACAGGGACCCCTGAGAAGCCCCTCTCAGAC CTTGGCCTCCTATCCTATCGAAGCTACTGGTCCCAGACCATCCTGGAGATCCTGATGGGGCTGAAG TCGGAGAGCGGGGAGAGGCCACAGATCACCATCAATGAGATTAGTGAAATCACCAGCATCAAGAAG GAGGATGTCATCTCCACTCTGCAGTACCTCAATCTCATCAACTACTACAAGGGCCAGTACATCCTC ACACTGTCAGAGGACATCGTGGATGGCCATGAGCGGGCCATGCTCAAGCGGCTCCTGCGGATCGAC TCCAAGTGTCTGCACTCCCCAAGGACTGGAGCAAGAGGGGGGAAGTGG

Figure 6 (continued)

>PTPK···

AGTAACTACATCAATGCTGCTCTTATGGACAGCTACAGGCAACCAGCTGCTTTCATCGTCACACAA
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GTGATGTTAAACGAAGTCGACTTGTCCCAGGGCTGCCCTCAGTACTGGCCAGAGGAAGGGATGCTA
CGATATGGCCCCATCCAAGTGGAATGTATGTCTTGTTCAATGGACTGTGATGTACAACCGGATT
TTTAGGATATGCAATCTAACAAGACCACAGGAAGGTTATCTGATGGTGCAACAGGTTTCAGTACCTA
GGATGGGCTTCTCATCGAGAAGTGCCTGGATCCAAAAGGTCATTCTTGAAACTGATACTTCAGGTG
GAAAAGTGGCAGGAGAATGCGAGGAAGGGGAAGGCCGGACGATTATCCACTGCCTAAATGGTGGC
GGGCGAAGTGGCATGTTCTGTGCTATAGGCATCGTTGTTGAAATGGTGAAACGGCAAAATGTTGTC
GATGTTTTCCATGCAGTAAAGACACTGAGGAACAGCAAGCCAAACATGGTGGAAGCCCCGGAGCAA
TACCGTTTCTGCTATGATGTAGGCTTTGGAGTACCTGGAATCATCT

Figure 6 (continued)

TTCTTCCCCATTAATTACGTGGAAGTGATCGTGCCTTTACCTCAG

>SH3GL3 GAAĞGAACTAAACTAGACGATGAATTTCTTGACATGGAAAGGAAAATAGATGTTACCAATAAAGTT cgaatgctgaacactgtgtcgaagatccgagggcaggtgaagaccacaggatacccgcagacggaa GGCTTGCTGGGGCACTCTATGCTGAAATACGGGAAGGCTCGGGGAAGACTCCACCTTTGGCAAT GCATTGATAGAAGTTGGTGAATCCATGAAGCTAATGGCTGAGGTGAAAGACTCTCTTGATATTAAT GTAAAGCAAACTTTTATTGATCCACTTCAGTTACTACAAGATAAAGATTTAAAAGAGATCGGGCAT CÁCCTGAÁAÁÁGCTGGAÁĞGCCGCCĞÖCTGGATTÁCGAŤTÁTÁAAAÁĞAÁÁCGÁGTÁGGTAÁGATA CCAGACGĂĂGAAGTCAGACAAGCGGTAGĂAAAATTTGAAGAGTCAAAGGAGTTGGCTGAAAGAAAGC atgittaactititagaaaatgatgtagaagtcagccagitegctgtgttcatagaggcagca TTAGACTÁTCACAGACAGTCCÁCAGAGÁTTCTCCÁGAGGAGCTGCÁGAGCAÁGCTACAGATCCCGÁATÁ TCAGCTGCATCCAGTGTCCCCAGACGACAAATACAAGGCCAGGCCTGTGAAAAGGACTTCTAGTGAG CTCAATGGAGTTTCCACCACCTCTGTAGTGAAGACGACAGGTTCTAACATTCCCATGGACCAGCCC TGCTCTCGTGGTCTCTATGACTTTGAGCCAGAAAACCAAGGAGAATTAGGATTTAAAGAAGGGGAC ATCATTACATTAACCAATCAAATAGATGAAAACTGGTATGAAGGAATGATACACGGAGAATCGGGA

PCT/EP2004/006617

CGGCCCCGCGCACAGTTGCGGCGGGAGAGCGGCGGGGCCGAGAGCGTGACTCGCCCGCTCCGCGCT GTGAAGACAGAGAATGACCACATCAACCTGAAGGTGGCCGGGCAGGACGGCTCCGTGGTGCAGTTC AAGATCAAGAGGCACACGCCGCTGAGCAÄGCTGATGAAGGCCTACTGCGAGAGGCAGGGCTTGTCA gagatggaggaggaggacaccatcgacgtgttccagcagcagacgggaggtgtgccagagaggcagc CTGGCAGGGCACAGTTTC

>SÜMO-3

CCCTCGTCCACCGCTGCCGCCTCCTTCTTCTGCCGCTCCTGGTGCTGCTTGTGTGTGCTCGTTTGGTG ccgacctgctaectcttttetcaagcgcacctgaggagactccgccetcgccatgccgacgaa AAGCCCAAGGAAGGAGTCAAGACTGAGAACAACGATCATATTAATTTGAAGGTGGCGGGCAGGAT GGTTCTGTGGGGGAGTTTAAGATTAAGAGAGCATACACCACTTAGTAAACTAATGAAACTATTGT GAACGÁCAGGGÄTTGTCAATGAGGCAGÁTCAGATTCCGÁTTTGACGGCAACCAÁTCAÁTGAAACA GACACÁCCTGCÁCAGTTGGÁAÁTGGAGGATGÁAGATACAATTGÁTGTTTCCAÁCAGCAGACGGGÁ GGTGTCTAC

AGCTCACCCTGAAGCGTCAGAGGATGGAGTCCGCGCTGGACCAGCTCAAGCAGTTCACCACCGTG ÐÖÖDÁÁDDÁÖÐĀŤÖÐTAÐÐÁÐÓÐÐÐAKGAÍTÐAÐDAÐ,TÁDÓÐDÁÚÍTÐÁÐÐÐÐÐÓÁÓAÐÐÐ TCCCTGATCCTGGCCGCACACACACACCCCCTTACCAGGGCCTGGTGGAGGAGGCGATTGCCTAT GGCCGGAAGCTGGGCGGCTCACAAGAGGACCAGATTAAAAATGCTATŤGÄTAAACTTTTTGTGTTG TTTGGAGCAGAAATACTAAAGAAGATTCCGGGCCGAGTATCCACAGAAGTAGACGCAAGGCTCTCC TTTGATAAGATGCGATGGTGGCCAGAGCCAGGCGGCTCATCGAGCTGTACAAGGAAGCTGGGATC GAGGAGCACCGCATCCACTGCAACATGACGTTACTCTTCTCCTTCGCCCAGGCTGTGGCCTGT GCCGÄGGCGGTGTGACCCTCATCTCCCCÄTTTGTTGGGCCCATCCTTGATTGGCATGTGGCAAAC AUCGACAAGAATCCTATGAGCCCCTGGAAGACCCTGGGGTAAAGAGTGTCACTAAAATCTACAAC TACTACAAGAGTTTAGCTACAAAACCATTGTCATGGGGGGCTCCTTCCGCAACACGGGCGAGATG AAAGCACTGGCCGGCTGTGACTTCCTCACCATCTCACCCAAGCTCCTGGGAGAGCTGCTGCAGGAC <u> ĎACĠĊĠAĠĊŦĠĠŢĠĊŢĠŢĠĊŤĊŢĊĄĠĊĠĀĠĠĠĠĊĊĊĄĠĊĊĠŢĠŔŢĊĊŢĠĠ</u> GCAGAGAATGGAAAG

TCCGGGAGCCAGTCCGCGCCACCGCCGCCGCCCAGCCATCGCCACCCTCCGCAGCCATGTCCACC AGGTCCGTGTCCTCGTCCTACCGCAGGATGTTCGGCGGCCCCGGGCACCGCGAGCCGGCCAGC TCCAGCCGGAGCTACGTGACTACGTCCACCCGCACCTACAGCCTGGGCAGCGCGCTGCGCCCCAGC ACCAGCCGCAGCCTCTACGCCTCGTCCCCGGGCGGCGTGTATGCCACGCGCTCCTCTCCCGTGCGC ATCAACACCGAGTTCAAGAACACCCGCACCAACGAGAAGGTGGAGCTGCAGGAGCTGAATGACCGC TTCGCCAACTACATCGACAAGGTGCGCTTCCTGGAGCAGCAGAATAAGATCCTGCTGGCCGAGCTC GAGCAGCTCAAGGGCCAAGGCAAGTCGCGCCTGGGGGACCTCTACGAGGAGGAGATGCGGGAGCTG CGCCGCAGGTGGACCAGCTAACCAACGACAAAGCCCGCGTCGAGGTGGAGCGCGACAACCTGGCC ACCCTGCAATCTTTCAGACAGGATGTTGACAATGCGTCTCTGGCACGTCTTGACCTTGAACGCAAA GTGGAATCTTTGCAAGAGAGAGTTGCCTTTTTGAAGAAACTCCACGAAGAGGAAATCCAGGAGCTG CAGGCTCAGATCCAGGAACAGCATGTCCAAATCGATGTGGATGTTTCCAAGCCTGACCTCACGGCT GCCCTGCGTGACGTACGTCAGCAATATGAAAGTGTGGCTGCCAAGAACCTGCAGGAGGCCAGAAGAA TGGTACAAATCCAAGTTTGCTGACCTCTCTGAGGCTGCCAACCGGAACAATGACGCCCTGCGCCAG GCAAAGCAGGAGTCCACTGAGTACCGGAGACAGGTGCAGTCCCTCACCTGTGAAGTGGATGCCCTT AAAGGAACCAATGAGTCCCTGGAACGCCAGATGCGTGAAATGGAAGAGAACTTTGCCGTTGAAGCT GCTAACTACCAAGACACTATTGGCCGCCTGCAGGATGAGATTCAGAATATGAAGGAGGAAATGGCT CGTCACCTTCGTGAATACCAAGACCTGCTCAATGTTAAGATGGCCCTTGACATTGAGATTGCCACC TACAGGAAGCTGCTGGAAGGCGAGGAGAGCAGGATTTCTCTGCCTCTTCCAAACTTTTCCTCCCTG AACCTGAGGGAAACTAATCTGGATTCACTCCCTCTGGTTGATACCCACTCAAAAAGGACACTTCTG ATTAAGACGGTTGAAACTAGAGATGGACAGGTTATCAACGAAACTTCTCAGCATCACGATGACCTT GAA

Figure 6 (continued)

>VIMc

>ZHX1

<u>GAATCTACAGAAGTTTCTTCTTCGGGAATATCTATCAGTAAAACTCCTATCATGAAAATGATGAAA</u> AATAAAGTGGAAAATAAACGGATTGCAGTTCATCATAACTCAGTTGAGGACGTTCCTGAAGAAAA GAGAATGAAATCAAACCAGACCGTGAAGAAATTGTAGAAAATCCAAGTTCTTCAGCTTCTGAATCT AATACAAGTACTTCCATTGTAAACAGAATACATCCAAGTACTGCCAGCACGGTAGTGACACCAGCA GCAGTTCTTCCTGGATTGGCACAGGTGATAACTGCTGTATCTGCTCAGCAGAATTCTAATTTGATT CCCAAAGTCTTAATCCCTGTTAATAGCATTCCCACCTACAATGCTGCATTGGATAACAATCCCCTT GCAAAATATACAGAGGAACAGATCAAGATATGGTTTTCAGCCCAACGTTTAAAACATGGTGTTAGT TGGACTCCCGAGGAAGTAGAGGAGGCAAGAAGGAAACAATTCAATGGAACAGTGCATACTGTACCT CAGACCATAACTGTTATTCCTACACACTTTCCACAGGGAGTAATGGTTTACCATCTATTTTACAG GTTACAGCACCTATAGCCTTGACAGTGGCAGGCGTTCCAAGTCAAAATAATATACAGAAAAGTCAG GTACCTGCTGCTCAGCCTACTGCAGAAACAAAGCCAGCAACAGCAGCAGTTCCAACTTCTCAAAGT GTCAAACATGAAACTGCATTGGTAAACCCTGATTCATTTGGCATTCGGGCAAAAAAAGACAAAAGAG CAACTGGCAGAATTAAAAGTTAGCTACCTTAAAAATCAGTTTCCCCATGATTCAGAAATTATCAGA CTTATGAAAATAACAGGCCTGACGAAAGGAGAGATTAAAAAATGGTTTAGTGACACAAGGTACAAC CAGAGAAATTCAAAGAGTAATCAGTGCTTACATCTCAACAATGATTCCTCTACCACCATTATTATA GACTCCAGTGATGAAACCACGGAATCCCCAACTGTTGGTACTGCACAGCCTAAGCAATCCTGGAAT CCTTTTCCTGACTTTACTCCCCAAAAGTTTAAAGAGAAAACTGCAGAGCAGCTTCGTGTCCTTCAG GCAAGTTTTCTCAACAGCTCTGTACTTACAGATGAAGAATTAAATAGGTTAAGGGCACAAACCAAA CTTACCAGAAGAGAATCGATGCTTGGTTTACAGAGAAGAAGAATCAAAAGCTTTAAAGGAAGAG AAAATGGAAATAGATGAAAGTAATGCAGGTAGTTCCAAAGAAGAAGCTGGAGAAACTTCTCCTGCA GATGAATCTGGTGCACCTAAGTCAGGGAGTACAGGCAAGATATGTAAAAAAACACCTGAGCAGCTG CACATGCTTAAGAGTGCATTTGTCCGGACACAGTGGCCATCACCAGAAGAGTATGACAAGTTGGCC AAAGAAAGCGGGCTTGCTAGAACAGACATAGTTAGTTGGTTTGGGGACACCCGTTATGCTTGGAAG $\verb"TCCCTTAGGAAAGAGGGAGAGGGAGACCCAAAGGACGGGGAAGAGGAAGACCGCGTGGGCGCCT"$ AGAGGAAGCAAAAGAATTAACAACTGGGACAGGGGACCATCACTCATAAAATTTAAAACTGGAACT GCAATACTTAAGGATTATTACCTGAAGCACAAGTTTCTTAATGAGCAAGACCTTGATGAACTTGTT AACAAATCACATATGGGCTATGAGCAGGTCAGAGAGTGGTTTGCAGAAAGACAGAGAAGATCAGAA GAAGAAGAAACAGATGATAGTGACACTTGGGAACCTCCACGACATGTGAAACGGAAGCTGTCTAAA TCAGATGAC

. Figure 6 (continued).

>ZNF33B

Figure 6 (continued)

Nucleotide sequence data (fasta format)

>GDF9

>GAPD

CCTGTTCGACAGTCAGCCGCATCTTCTTTTGCGTCGCCAGCCGAGCCACATCGCTCAGACACCATGGGGAAGGTG
AAGGTCGGAGTCAACGGATTTGGTCGTATTGGGCGCCTGGTCACCAGGGCTGCTTTTAACTCTGGTAAAGTGGAT
ATTGTTGCCATCAATGACCCCTTCATTGACCTCAACTACATGGTTTACATGTTCCAATATGATTCCACCCATGGC
AAATTCCATGGCACCGTCAAGGCTGAGAACGGGAAGCTTGTCATCAATGGAAATCCCATCACCATCTTCCAGGAG
CGAGATCCCTCCAAAATCAAGTGGGGCGATGCTGGCGCTGAGTACGTCGTGGAGTCCACTGGCGTCTTCACCACC
ATGGAGAAGGCTGGGGCTCATTTGCAGGGGGGGAGCCAAAAGGGTCATCATCTCTGCCCCCTCTGCTGATGCCCCC
ATGTTCGTCATGGGTGTGAACCATGACAAGTATGACAACAGCCTCAAGATCATCAGCAATGCCTCCTGCACCACC
AACTGCTTAGCACCCCTGGCCAAGGTCATCCATGACAACTTTTGGTATCGTGGAAGGACTCATGACCACAGTCCAT
GCCATCACTGCCACCCAGAAGACTGTGGATGGCCCCTCCGGGAAACTGTGGCGTGATGGCCGCGGGGCTCTCCAG
AACATCATCCCTGCCTCTACTGGCGCTGCCAACGTGTGGGCAAGGTCATCCCTGAGCTGAACAGGAAGCTCACT
GGCATGGCCTTCCGTGTCCCCACTGCCAACGTGTCAGTGGTGGACCTGACCTGCCGTCTAGAAAAAACCTGCCAAA
TATGATGACATCAAGAAGGTGGTGAAGCAGGCGTCGGAGGGCCCCCTCAAGGGCATCCTGGGCTACACTGAGCAC
CAGGTGGTCTCCTCTGACTTCAACAGCGACACCCACTCCTCCACCTTTGACGACACAGGGTTGGCCTCAAC
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CACATGGCCTCCAAGGAG

>MOV34

Protein sequence data (fasta format)

->GDE9

 ${\tt HSWYSLHYKRRPSQGPDQERSLSAYPVGEEAAEDGRSSHHRHRRGQETVSSELKKPLGPASFNLSEYFRQFLLPQ\\ {\tt NECELHDFRLSFSQLKWDNWIVAPHRYNPRYCKGDCPRAVGHRYGSPVHTMVQNIIYEKLDSSVPRPSCVPAKYS\\ {\tt PLSVLTIEPDGSIAYKEYEDMIATKCTCR}.$

>GAPD

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>MOV34

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Figure 6 (continued)

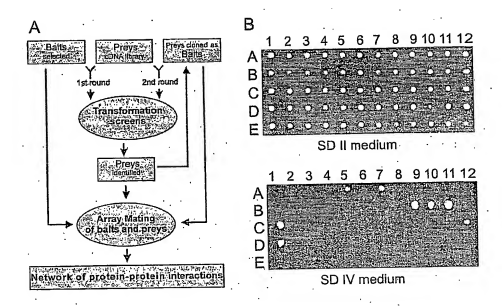
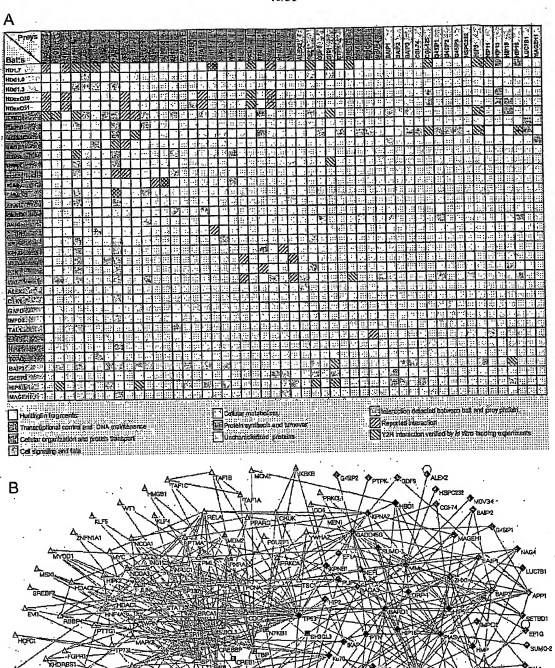


Figure 7

→ PFN2 → HP28

Figure 8

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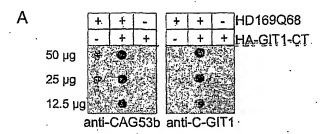
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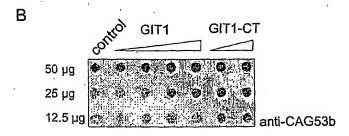
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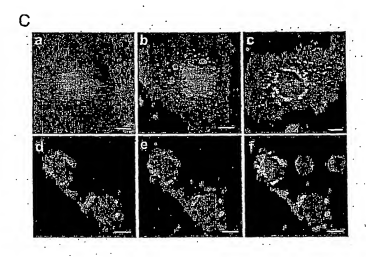
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GIT1	HD510Q17	
HZFH	HD510Q17	
HIP11	HD510Q17	
HIP1	HD510Q17	5.000 mg
IKAP	HD510Q17	
Ku70.	HD510Q17	
PFN2	HD510Q17	
PIASy	HD510Q17	200
HIP5	HD510Q68	
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HIP5	GIT1	
HZFH	GIT1 /	
HIP15	BAIP3	
GIT1	BAIP3	
HIP5	HBO1	
BARD1	HBO1	
BARD1	HIP5	
GADD45G	HIP5	
sumo-3	PIASy	
hADA3	PIASy	
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GADD45G	PLIP	
BARD1	CA150	
BARD1	HZFH	
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HIP5	FEZ1	
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Figure 9







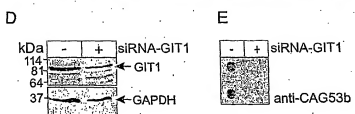
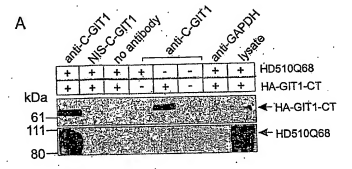
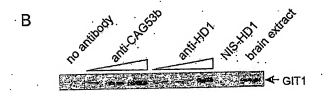


Figure 10





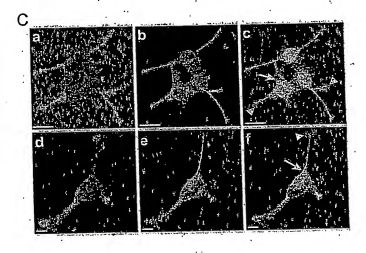


Figure 11

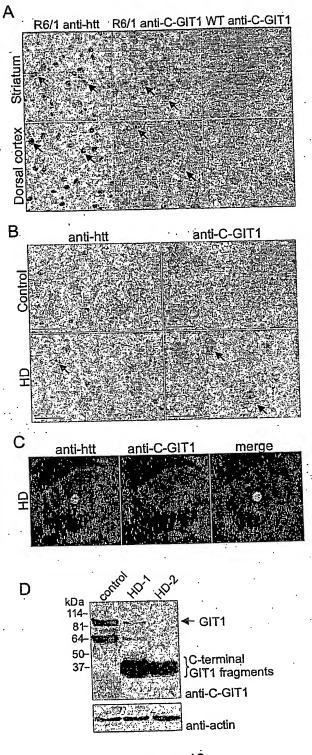
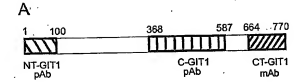


Figure 12



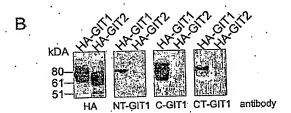


Figure 13

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